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WU and KI polyomavirus present in the respiratory tract of children, but not in immunocompetent adults

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Abstract

Background: Recently, two new polyomaviruses (PyV), termed WUPyV and KIPyV, were identified in respiratory tract specimens from children with acute respiratory tract infections (ARTIs). However, their roles in the disease have not been determined.

Objectives: To determine the prevalence of WUPyV and KIPyV in the Chinese population suffering from ARTIs in Beijing, China, and to examine their possible role in causing disease.

Study design: Nasopharyngeal aspirates, nasal swabs and throat swabs were collected from 415 children and 297 immunocompetent adults with lower ARTIs. The specimens were screened by polymerase chain reaction for the presence of WUPyV, KIPyV, and other common respiratory pathogens.

Results: Although none of the adults sampled were positive for either virus, WUPyV in 10 (2.4%) children and KIPyV was detected in 2 (0.5%) of the children sampled, respectively. Eleven of the positive cases were co-detected with either rhinovirus (6/11), respiratory syncytial virus (4/11), parainfluenzavirus virus (3/11) or Mycoplasma pneumoniae (2/11). Phylogenetic analysis of the WUPyV and KIPyV isolates showed that the nucleotide sequences were homologous to those of previously reported strains.

Conclusions: The presence of WUPyV and KIPyV in samples from children but not from immunocompetent adults suffering from LARTIs suggests that these viruses primarily infect the young population. Co-detection of additional respiratory pathogens in most of the specimens containing either WUPyV or KIPyV suggests that these viruses do not cause disease independently.

1. Introduction

Two new polyomaviruses (PyV), termed WUPyV and KIPyV, have been identified in respiratory tract specimens from children with acute respiratory tract infections (ARTIs), suggesting a potential role for polyomaviruses in human ARTIs. Full-length genome sequencing demonstrated that WUPyV and KIPyV are genetically similar and together form a new subfamily within the Polyomaviridae. Previous studies have examined the prevalence of these two novel viruses within different regions, such as Australia, Korea, Canada, USA, Germany, and Thailand, and their association with the population suffering from ARTIs. However, the clinical significance of WUPyV and KIPyV remains to be established. In order to further investigate the association between viral infection and the manifestation of ARTIs, we examined the prevalence, as well as clinical and molecular features, of
WuPyV and KIPyV infection among children and adults with lower ARTIs (LARTIs) in Beijing, China.

2. Methods

Nasopharyngeal aspirates (NPAs) were collected, upon admission, from 415 hospitalized children with LARTIs (262 males and 153 females) at the Beijing Children’s Hospital between March and December 2007. Children ranged in age from 1 month to 14 years, with a mean age of 2.6 years and a median age of 8.2 months.

Nasal and throat swabs were collected from 297 immunocompetent adult patients (148 males and 149 females) when they were diagnosed with LARTIs at the Outpatient Clinic of the Peking Medical College Hospital between May 2005 and December 2007. Of the 297 adults sampled, 74 were identified and sampled during the same period as that for the pediatric study described above. For each adult, both nasal and throat swabs were collected simultaneously and pooled into one tube containing virus transport medium. Adults ranged in age from 15 years to 97 years, with a mean age of 47.2 years and a median age of 43 years.

Viral nucleic acids were extracted from each specimen using the NucliSens easyMAC™ system and the NucliSens Isolation Reagents (bioMérieux). All specimens were tested for the presence of common respiratory viruses including parainfluenza viruses (PIV) 1–4, human rhinovirus (HRV), enterovirus, influenza viruses, respiratory syncytial virus (RSV), human coronaviruses (229E, OC43, NL63 and HKU1), human metapneumovirus, adenovirus, and human bocavirus using several molecular tests, as previously described.10–15 The presence of Mycoplasma pneumoniae (MP), was established using MP-specific IgM antibody and the gelatin particle agglutination test kit (SERODIA-MYCO II, Fujirebio Inc., Japan).

WuPyV was detected by PCR analysis using primers targeting its VP2 gene, generating a 250-bp amplicon.1 In addition, to exclude the false-negative results, the WuPyV samples negative for VP2 were retested using primers targeting the large T antigen (LTAg) gene.1 KIPyV was detected by nested PCR using primers targeting its VP1 gene,2 amplifying a 207-bp fragment. BKPyV and JCPyV were detected by PCR analysis as described previously before.1 KIPyV was detected in two pediatric patients (0.5%) aged 3 months and 3.5 years. Of the two KIPyV-positive cases, one was positive for KIPyV alone, while the other was positive for both KIPyV and PIV2.

None of the 297 samples from adults tested positive for WuPyV or KIPyV, and none of the pediatric or adult patient samples tested positive for the presence of BK or JC PyVs.

The 12 children who tested positive for either WuPyV or KIPyV suffered a wide range of respiratory diseases including bronchopneumonia (7/12), peribronchitis (2/12), and pneumonia (3/12) (Table 1). In addition, five had underlying serious medical conditions. Notably, patient BCH-370A, who was hospitalized due to

### Table 1

| Patient no. | Age/sex | Date of collection | Symptoms/health deficiencies or sign | Clinical diagnosis | Time in hospital (days) | Pathogens detected |
|-------------|---------|--------------------|--------------------------------------|--------------------|------------------------|-------------------|
| BCH-151A    | 3.5 years/F | August 14          | Cough, fever, low O 2 sat              | Peribronchitis      | –                      | KIPyV + PIV2      |
| BCH-370A    | 3.6 months/M | December 15        | Cough, gasping, premature (33 w), patent foramen ovale, newborn wet lung, newborn aspiration pneumonitis, bronchopulmonary dysplasia cough, vomit, diarrhea, fever | Pneumonia Congestive heart failure | 11                   | KIPyV             |
| BCH-004A    | 8.5 months/M | March 27           | Cough, gasping, premature (33 w), patent foramen ovale, newborn wet lung, newborn aspiration pneumonitis, bronchopulmonary dysplasia cough, vomit, diarrhea, fever | Bronchopneumonia Myocardial damage | 9                    | WuPyV + RSV       |
| BCH-34A     | 2.2 months/F | April 12           | Cough, gasping, fever/patent ductus arteriosus | Bronchopneumonia Myocardial damage | –                     | WuPyV + HRV       |
| BCH-123A    | 2.7 mon/M | July 30            | Cough, gasping, fever/patent ductus arteriosus | Bronchopneumonia Myocardial damage | 6                    | WuPyV + PIV3 + PIV4 |
| BCH-172A    | 4.3 y/M | September 6          | Cough, gasping, fever/patent ductus arteriosus | Bronchopneumonia Myocardial damage | 12                   | WuPyV + PIV4      |
| BCH-200A    | 8 months/M | October 15          | Cough, gasping, fever/patent ductus arteriosus | Peribronchitis      | –                     | WuPyV + HRV       |
| BCH-235A    | 2.9 years/M | October 29         | Cough, gasping, fever/patent ductus arteriosus | Bronchopneumonia Myocardial damage | 13                   | WuPyV + HRV       |
| BCH-259A    | 2.7 years/M | November 8          | Cough, fever                           | Pneumonia Pleural effusion | 14                   | WuPyV + HRV       |
| BCH-312A    | 1.2 months/M | November 22        | Cough, fever                           | Bronchopneumonia Myocardial damage | 15                   | WuPyV + HRV + RSV  |
| BCH-313A    | 11 months/M | November 25         | Cough, vomit, fever/metabolic acidosis | Bronchopneumonia Cardiac insufficiency | 7                    | WuPyV + HRV + RSV  |
| BCH-404A    | 7.4 years/F | December 24         | Cough, nausea, vomit, fever            | Pneumonia           | 7                     | WuPyV + RSV + MP   |

KIPyV, KI polyomavirus; PIV, parainfluenzavirus; WuPyV, Wu polyomavirus; RSV, respiratory syncytial virus; HRV, Rhinovirus; MP, Mycoplasma pneumoniae.
pneumonia and tested positive for KIPyV alone, had severe health problems including prematurity (born at 33 weeks), patent foramen ovale, newborn wet lung, newborn aspiration pneumonitis, and bronchopulmonary dysplasia. Each of the WUPyV- or KIPyV-positive patients eventually recovered.

Phylogenetic analysis revealed a very high level of identity between the 10 WUPyV isolates (GenBank accession numbers: EU597008, EU754876–83, EU784672), as well as between the 2 KIPyV isolates (GenBank accession numbers: EU754874–75). The nucleotide sequences of the VP2 gene from the WUPyV strains found showed 98–100% homology with previously described strains. The VP1 gene sequences from the KIPyV isolates showed 99–100% homology with the reference strains (Fig. 1). Despite the limited sequences analyzed, the minimal variations suggest the stability of the virus genomes and the global similarity of the epidemic strains.

Fig. 1. Phylogenetic analysis of nucleotide sequences of VP2 of WUPyV (a) and VP1 of KIPyV (b) isolates (the isolates identified during this study are shown in italics). The tree was built with the MEGA4.0 software by using distance method and the neighbor-joining algorithm with Kimura 2 parameters. Strains isolated in Beijing are indicated by a specific identification code (BCH) followed by the patient number. The analysis included WUPyV reference sequences from GenBank including B0, B12, B3, B14, B5, B8, B7, B28, B2, KR-M-2217, CLFF, KR-M-3293, KR-M-2338 (GenBank accession numbers: EF444550, EF444557, EF444558, EF444562, EF444563, EF444579, EF444580, EF444590, EF444591, EF655819, EU296475, EU041606, EF655820, respectively), and KIPyV reference sequences including LZ65, LZ285, KR-M-3209, KR-M-3292, Stockholm 60, Brisbane 002, Stockholm 380, Mpt2 (GenBank accession numbers: EU309498, EU309496, EF639289, EU041609, NC_009238, EF520288, EF127908, AM849809).
4. Discussion

Our investigation provides valuable insights into the population that is susceptible to infection with these viruses. Consistent with previous reports, our finding indicates that the majority (8/10) of WUPyV-positive samples originated from children less than 3 years old. However, because the positive sample number is low, further investigation with larger positive patient pools is required to accurately estimate the age distribution of WUPyV and KIPyV infections.

In an effort to further study the clinical relevance of WUPyV and KIPyV infection in LARTIs, we assayed for the presence of these viruses in immunocompetent adults diagnosed with LARTIs. Significantly, no positive cases were detected among the population sampled. Although prior studies have reported the presence of WUPyV and KIPyV in adults, the populations sampled in these studies corresponded to immunocompromised patients suffering from WUPyV and KIPyV in adults only occurs when they are immunosuppressed or have other significant health deficiencies. However, before making this conclusion, we must consider the limitations associated with particular specimen types on viral detection. In our study, adult specimens were collected using throat and nasal swabs. As the viral load in such specimens is usually lower than in NPAs, it is possible that the prevalence of WUPyV and KIPyV in adults has been considerably underestimated. Future studies focusing on the prevalence of WUPyV and KIPyV in adults should therefore assay lower respiratory tract samples, such as bronchoalveolar lavage fluid (BALF).

All but one of the WUPyV- or KIPyV-positive pediatric patients were co-infected with other respiratory pathogens, suggesting that WUPyV and KIPyV, may not directly cause disease, but rather act by one, or a combination, of the following: (1) as opportunistic pathogens in LARTIs; (2) colonizing the respiratory tract without causing any disease; (3) a part of the endogenous viral flora, which could be reactivated by other viral infections; or (4) playing a role in causing severe diseases in the presence of other pathogens. One pediatric patient was positive for KIPyV alone. However, the possibility that this patient was also infected with other pathogens cannot be excluded as we assessed only for the presence of common respiratory viruses and MP during this study.

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