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Correlation between bocavirus infection and humoral response, and co-infection with other respiratory viruses in children with acute respiratory infection

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1. Background

In 2005, Human bocavirus (HBoV), a novel member in the genus Parvovirus, family Parvoviridae, was first identified in children with lower respiratory tract infection (LRTI). Since then, the detection of HBoV in acute respiratory illness was reported worldwide, with prevalence rates between 1.5% and 19%. Among parvoviruses, HBoV has the closest amino acid identity with bovine parvovirus (42%) and canine minute virus (43%). HBoV is a single-stranded DNA virus with a genome of around 5 kb, encoding 4 genes (NS1, NP1, VP1, and VP2). The majority of genetic variations occurred in the capsid VP1/VP2 genes, allowing a classification into two genotypes, ST1 and ST2.

HBoV was detected in various samples, including nasopharyngeal aspirate, serum, feces, and urine. HBoV detection was significantly higher in patients with symptoms of respiratory tract or gastroenteritis than in asymptomatic individuals. In addition, higher HBoV prevalence was observed in children aged from 6 months to 2 years, compared to older individuals. Co-infection with other respiratory viruses was commonly observed in HBoV-positive patients with respiratory tract symptoms. Thus, whether HBoV is a major cause of respiratory disease remains questionable.

Recently, several studies on the seroepidemiology of HBoV were reported by immunofluorescence assay, Western blot, or ELISA. Anti-HBoV IgG was detected in a large fraction of the population, especially in older children and adults, while IgM was found predominantly in HBoV-positive patients.

Only few studies have undertaken a comprehensive analysis of clinical, virological, and serological aspects in HBoV infection. The present study aimed at better characterizing the virological and serological profiles of HBoV in children with acute respiratory infection (ARI) in samples collected during a 2-year period in one hospital in Shanghai, China. Co-infected respiratory viruses were analyzed, and serological diagnosis using a highly antigenic fragment of VP2 was developed. In addition, persistent HBoV shedding was assessed in nasal, throat, serum, and urine specimens.
2. Materials and methods

2.1. Patients and specimens

From October 2006 to September 2008, 817 samples were collected from children aged between 6 months and 9 years, who were diagnosed with an ARI with accompanying fever. All specimens were collected at the Shanghai Nanxiang Hospital, with informed consent from the children’s parents. Standardized protocols accepted by the ethical committee of the hospital and reviewed by the medical committee of Institut Pasteur in Paris were used to record the patients’ clinical symptoms and medical history, and for specimen collection. Nasal and throat swab samples were collected and stored in viral transport medium, and blood samples were collected in tubes containing 3.2% sodium citrate. The doctor who was involved in the study and did the sampling remained the same throughout the study. All samples were aliquoted at arrival who was involved in the study and did the sampling remained the same throughout the study. All samples were aliquoted at arrival and stored at −80 °C. Twelve patients who were diagnosed positive for HBoV were recalled between 8 and 31 days after the initial visit, and nasal swabs, throat swabs, blood, as well as urine samples were collected.

2.2. Quantitative PCR for detection of HBoV in human sample extracts

Total RNA and DNA were extracted from 140 μl of each specimen by QIAamp Viral RNA Mini Kit (QIAGEN, Hilden, Germany), according to the manufacturer’s instructions. Nucleic acids were then eluted in 50 μl RNase-free water and stored at −80 °C until use. The primers and the probe used to detect HBoV were targeted to the NP1 gene, as published previously (Table 1).26 The real-time PCR assays were performed using Takara’s Premix Ex TaqTM Perfect Real Time (Takara Biotechnology, Dalian, China) in 20 μl reaction mixture, containing 1 μl of extracted sample, 0.2 μM of each of the forward and reverse primers, and 0.2 μM of probe. PCR amplification included a step at 95 °C for 10 s followed by 40 cycles of 5 s at 95 °C and 30 s at 60 °C. The reactions were run and analyzed in an ABI PRISM 7900HT (Applied Biosystems, Foster City, CA, USA). The standard curve was obtained using serial dilutions of pskHBoV containing the whole genome of HBoV ST2 genotype (a kind gift from Dr Jianming Qiu, University of Missouri-Columbia, USA). The housekeeping gene GAPDH was detected as the internal control to monitor the quality of the specimens and to normalize HBoV DNA in clinical samples for comparative analysis.

2.3. Conventional RT-PCR for HBoV and other respiratory viruses

HBoV-positive specimens were also examined with conventional RT-PCR, using the One-Step RT-PCR kit (QIAGEN) that targets the NS1 gene, as described previously.13 Multiplex RT-PCR, described previously, was used to diagnose other respiratory viruses in HBoV-positive samples,27 including respiratory syncytial virus (RSV), human rhinovirus (HRV), influenza A virus (IAV), influenza B virus (IBV), human metapneumovirus (HMPV), adenovirus (ADV), human coronaviruses (HCoV) 229E, OC43, HKU-1, and NL63, and parainfluenza viruses (PIV)1–4. Briefly, 2.5 μl of the sample extract and 0.2 μM of primers were added to the 25 μl reaction mixture. The RT reaction was carried out at 50 °C for 30 min, followed by PCR, including a step at 95 °C for 15 min and 40 cycles of 30 s at 94 °C, 30 s at 55 °C, 30 s at 72 °C and the final extension at 72 °C for 10 min. The products were run in a 1% agarose gel and visualized by UV light.

2.4. Sequencing of VP1 and phylogenetic analysis

The entire VP1 gene was amplified with primer pairs (Table 1) using QIAGEN’s One-Step RT-PCR kit (QIAGEN). The reaction mixture included 5 μl RNA, 0.4 μM of each primer, 400 μM of each dNTP, and 2 μl of enzyme mix in a final volume of 50 μl. Amplification included 45 min at 50 °C, 15 min at 94 °C, 8 cycles of 1st round amplification (30 s at 94 °C, 45 s at 55 °C, 2.5 min at 72 °C), followed by 35 cycles of 2nd round amplification (30 s at 94 °C, 45 s at 50 °C, 2.5 min at 72 °C), and final extension of 10 min at 72 °C. Products of 2016 bp were visualized in an agarose gel under UV light. The fragments were extracted by QIAquick Gel Extraction Kit (QIAGEN) and cloned into pMD18T (Takara Biotechnology). At least four colonies were picked from each sample and sent for sequencing to Biosune Carlsbad, CA, USA). The VP1 gene of HBoV strains collected during 2005–2009 available in Genbank were also downloaded and phylogenetic analysis was conducted with MEGA version 3.1 software. The phylogenetic tree was drawn by the Neighbor-joining method with bootstraps of 1000 replications.

2.5. Expression of VP2 in the baculovirus–insect cell system

The recombinant bacmid pFastBac1 containing HBoV VP2 gene (kindly provided by Dr Jianming Qiu) was transfected into Spodoptera frugiperda Sf21 cells to recover the recombinant baculovirus. High Five cells were infected with the virus at a multiplicity of infection of five plaque-forming units/cell and cells were harvested 3 days later by centrifugation. The cell pellet was resuspended in 25 mM NaHCO3 at a density of 2 × 106 cells/ml and left on ice for 30 min until the cells were completely lysed. Cell debris was removed by centrifugation at 13,000 rpm. The proteins were further precipitated in 20% ammonium sulfate, and resuspended in PBS.

2.6. Recombinant protein production

Four regions of VP2 were amplified by PCR from the pskHBoV clone. The forward and reverse primers (Table 1) contained Hind III and Xho I restriction sites, respectively. After digestion with

| Gene                      | Amplicon                  | Forward primer       | Reverse primer       | Size (bp) | Reference |
|---------------------------|---------------------------|----------------------|----------------------|-----------|-----------|
| NP1 for real-time PCR     | NP-1 Probe                | 5′-AGAGGCTGGGCTATATCA-3′ | 5′-CACCCTGGTCAGGTCTTGGAA-3′ | 81        | 26        |
| NS1 for conventional RT-PCR | NS-1                      | 5′-TATGGCCAAAGGATCGTCAAAG-3′ | 5′-GCCCGTGAAATGAGACAGC-3′ | 245       | 13        |
| VP1 for sequence          | VP1                       | 5′-CCACTAGTATGGCTTCAAATAGGACAGC-3′ | 5′-CTCTGGATTAAACATCTTTATATGGTGTTG-3′ | 2016      |           |
| VP2                       | VP2-P1                    | 5′-CCGCGTGGATATACAAACAGGAACTGGAACCG-3′ | 5′-CCGCGTGGATATACAAACAGGAACTGGAACCG-3′ | 480       | 13        |
| VP2                       | VP2-P2                    | 5′-CCGCGTGGATATACAAACAGGAACTGGAACCG-3′ | 5′-CCGCGTGGATATACAAACAGGAACTGGAACCG-3′ | 465       | 11        |
| VP2                       | VP2-P3                    | 5′-CCGCGTGGATATACAAACAGGAACTGGAACCG-3′ | 5′-CCGCGTGGATATACAAACAGGAACTGGAACCG-3′ | 474       | 11        |

Table 1

HBoV primers and probe used in viral genome amplification.
the restriction enzymes, the amplicons were cloned into pET30a containing 6xHis tag at the 5’ terminal. The recombinant proteins were expressed in *E. coli* BL21(DE3) host strains by induction with IPTG for 3 h. Cells were collected by centrifugation and the pellets were resuspended in lysis buffer and lysed by sonication on ice. The recombinant proteins were further purified by Ni-NTA agarose (Invitrogen) under denaturing conditions, following the manufacturer’s instructions. The protein needed for ELISA was further dialyzed into PBS.

### 2.7. Immunoblotting

Fifteen micrograms of each protein were loaded and separated on a 12.5% SDS-PAGE gel, and then transferred onto nitrocellulose membranes. After blocking for 2 h in 5% nonfat dry milk in TBS-T (0.5% Tween-20 in TBS buffer) blocking buffer, the membranes were incubated for 1 h in patient serum samples diluted in blocking buffer (dilution 1:50 for IgM test, 1:100 for IgG test). After incubation, the membranes were washed four times in TBS-T, and incubated in anti-human IgG or IgM conjugated with horseradish peroxidase (HRP) (Bethyl Laboratories, Montgomery, TX, USA). After 1 h, the membranes were washed four times in TBS-T buffer and 3,3’-diaminobenzidine and H$_2$O$_2$ were added to reveal the peroxidase activity.

### 2.8. ELISA

ELISA 96-well plates (Nunc™, Roskilde, Denmark) were coated with 200 ng/well of purified proteins in coating buffer (0.05 M carbonate/bicarbonate, pH 9.6) overnight at 4°C. After washing with PBS-T (0.05% Tween-20), the wells were saturated in 5% dry milk in PBS-T at 37°C for 2 h. Serum samples diluted in 1% dry milk in PBS-T were added and incubated for 1 h at 37°C. After washing, HRP-conjugated anti-human IgG (1:10,000 diluted in dry milk in PBS-T) was added for 1 h at 37°C. Plates were washed five times with PBS-T, and OPD (o-Phenylenediamine Dihydrochloride) (Sigma, St. Louis, MO, USA) was used as a substrate. After 20 min of incubation, the absorbance at OD$_{450}$ was read. All samples were measured in triplicate, and the average value was calculated for each serum sample. A purified heptapeptide fragment of SARS-CoV spike protein produced in *E. coli* was used as a negative antigen (the corresponding expression plasmid pet30a-SARS spike was a kind gift of Dr. YongJin Wang, East Normal University, Shanghai, China). The threshold for positivity was calculated as twice the mean OD$_{450}$ value obtained when sera were tested against the SARS spike protein fragment.

### 2.9. Statistical analysis

The statistical analysis was performed via SAS 6.12 software (SAS Institute Inc., Cary, NC, USA). The *p* value was measured by two-sided analysis, and a *p* value less than 0.05 was considered to be statistically significant. Comparisons between different variations were evaluated by chi-squared or Fisher’s test.

### 3. Results

#### 3.1. Viral screening from pediatric patients

During a 2-year period, a total of 817 pairs of nose and throat swabs were obtained from pediatric patients (male: 531, female: 286) with an age range from 6 months to 9 years. To detect HBoV, we first used conventional RT-PCR and then real-time PCR, targeting conserved NS1 and NP1 gene sequences, respectively. Samples with discrepant results between negative RT-PCR and positive real-time PCR were retested by increasing the amount of nucleic acid extract in conventional RT-PCR and by duplicating real-time PCR. Results finally gave 100% concordance between the two techniques. Ninety-six children (11.8%) were positive for HBoV detected in 96 throat swabs and 78 in nose swabs. Among 103 patients with virus co-infection, 49 (47.6%) were found infected with HBoV (data not shown). Forty two samples contained another virus, six samples contained 3 viruses, and one contained 4 viruses (Table 2).

The HBoV loads detected in nose and throat swabs ranged from <500 to 10$^6$ DNA copies/ml, and those in serum ranged from <500 to 10$^9$ copies/ml. Thirteen (59.1%) of 22 patients showing high copies (>10$^6$ copies/ml) of HBoV in throat swabs were also positive in their serum, whereas only 8 (14.0%) out of 57 patients showing low copies of HBoV were positive in their serum (*p*= 0.00012). Forty patients (62.5%) among 64 with a low level of HBoV copies in their throat were co-infected with another virus, whereas only nine (28.1%) from 32 with high copies of HBoV were co-infected (*p*= 0.00022).

#### 3.2. Clinical characteristics for HBoV-positive patients

HBoV was detected in 17.1% of children younger than 1 year, 14.3% in 1–2-year-old children, 8.2% in 2–4-year-olds, and 11.2% in older children. The gender ratio of HBoV-positive patients was 55% male and 45% female.

The main clinical symptoms in HBoV-positive patients involved fever (100%), cough (74.7%), and wheezing (20.9%). No specific symptom was prevalent in the HBoV-positive patients (Table 3). The number of HBoV RT-PCR-positive patients with bronchitis, however, was low (5.4%) when compared to the high prevalence of this disease (Table 3).

#### 3.3. Phylogenetic analysis of VP1 genes of HBoV

The VP1 gene of HBoV from 17 samples containing enough DNA/RNA material was amplified and sequenced. The nucleotide identity among different strains was over 99.3%. All strains tested in this study clustered with European, Asian and American strains into genotype ST2 (Fig. 1).

### Table 2

| Two virus infection (n) | Three virus infection (6 cases) | Four virus infection (1 case) |
|------------------------|--------------------------------|-------------------------------|
| RSV (10)               | PIV1/HRV                       | RSV/HRV/IBV                   |
| HRV (4)                |                                |                               |
| IAV/IBV (5/7)          | RSV/HMPV                       |                               |
| HMPV (3)               | RSV/229E                       |                               |
| ADV (5)                | PIV4/HBV                       |                               |
| HCoV-NL63 (1)         |                               |                               |
| PIV1/3 (4/3)          | RSV/ADV                        |                               |

RSV: Respiratory syncytial virus; HRV: human rhinovirus; IAV: influenza A virus; IBV: influenza B virus; HMPV: human metapneumovirus; ADV: adenoavirus; HCoV: human coronaviruses (including 229E, NL63); PIV1–4: parainfluenza virus 1–4.

### Table 3

| Clinical diagnosis | Total samples | HBoV-positive | Percentage of HBoV-positive/total samples |
|--------------------|---------------|---------------|-----------------------------------------|
| Acute laryngitis   | 58            | 11            | 19.0%                                   |
| Bronchitis         | 479           | 26            | 5.4%                                    |
| Bronchiolitis      | 51            | 11            | 21.6%                                   |
| Bronchiopneumonia  | 187           | 39            | 20.9%                                   |
| Asthma             | 42            | 9             | 21.4%                                   |
| Total              | 817           | 96            | 11.8%                                   |
3.4. Analysis of antigenic regions in VP2 of HBoV immunoreactive with patient sera

The VP2 capsid protein and four fragments of VP2 (designated P1 to P4) were first tested in a panel of serum samples by Western blots to detect HBoV-specific IgG and IgM antibodies. Eight of the samples contained high HBoV load, six contained low viral load, and six contained no HBoV. VP2 and its P3 fragment were the most immunoreactive with both IgG and IgM, and the P1 fragment was reactive mostly with IgM (Fig. 2). Therefore, the 79 available serum samples from HBoV RT-PCR-positive patients were tested in Western blots against VP2 protein and P1 and P3 polypeptides.
to detect IgM and IgG. In total, 40 (50.6%) sera were IgG-positive, and 44 (55.7%) were IgM-positive for HBoV antigens. Nineteen of 21 HBoV viremic patients had antibodies, with 15 (71.4%) IgM-positive and 11 (52.3%) IgG-positive. Among 22 children infected with high HBoV DNA levels (>10⁴ copies/ml) in nasal or throat swabs, 16 (72.7%) were IgM-positive and 13 (59.1%) were IgG-positive. In 36 single-infected specimens, 26 (69.4%) were IgM-positive, compared with 18 (41.9%) in 43 co-infected samples (p = 0.012) (Table 4).

3.5. Analysis of anti-HBoV IgG antibodies by ELISA using VP2-derived P3 fragment

The sera of patients were tested in ELISA using a purified P3 antigenic fragment derived from the VP2 protein, and SARS-HCoV (HR2 spike protein fragment) as a negative control. At a serum dilution of 1:100, the mean OD₄₅₀ value for negative controls using the SARS-HCoV spike HR2 fragment was 0.165. The assay was further confirmed by Western blots and the ELISA values were considered either negative (OD₄₅₀ < 0.33 corresponding to twice the mean value of the negative control), equivocal (OD₄₅₀ = 0.33–0.42), or positive (OD₄₅₀ > 0.42).

In all, 713 serum samples were available to test HBoV-specific IgG antibodies. 314 samples (44.0%) were seropositive, and 95 (13.3%) were equivocal. The number of IgG-positive individuals increased with their age, with a statistical significance for age distribution (p < 0.0001) (Fig. 3). Fewer children younger than 2 years old had anti-HBoV IgG (41.2%) than those older than 4 years old (67.93%) (p < 0.0001). In addition, 17 out of 29 sera showing an OD₄₅₀ value above 0.85 were from children older than 4 years.

3.6. Persistent shedding of HBoV

Twelve HBoV-positive patients were called back around 2 weeks after the first sampling. The mean interval of specimen collection was 16 days (days 8–31). None of these children showed any symptoms of ARI, and obviously recovered from their previous infection. However, HBoV could still be detected in nose and throat swabs, serum or urine (Table 5) from 10 children with a viral load ranging from less than 10² to 10⁴ copies/ml. In all four specimens, the virus loads were lower or similar in the second samples compared to the first samples, and two samples were found to be free of HBoV. No urine samples were collected during the first sampling, and three urine specimens were positive for HBoV in return-visit samples. In one case, virus genes were still detected in nose/throat swabs as well as in urine collected 31 days apart. Seven serum samples showed a seroconversion for IgG and three for IgM. All but two samples contained IgM in return samples. An anti-HBoV IgG antibody response was detected in all but two patients showing low virus titer during the second visit. Co-infecting viruses found in four children at the first visit were absent at the second visit but HRV or ADV were newly detected in three children showing low levels of persistent HBoV (Table 5).

4. Discussion

In this study, the prevalence rate of HBoV-positive non-hospitalized infants with ARI was as high as 11.8%, and infants less than 2-year-old were more frequently positive than older ones. This is similar to that reported by other studies, 14–16 All the HBoV strains clustered in the ST2 genotype, as previously identified in

### Table 4

Antibody responses in sera (via Western blot) compared with viral loads in HBoV-positive samples.

| Viral load/Ab response            | Total samples | IgM no. (%) | IgG no. (%) | Antibodies no. (%) |
|----------------------------------|---------------|-------------|-------------|--------------------|
| Viremia                          | 21            | 15 (71.4%)  | 11 (52.4%)  | 19 (90.5%)         |
| With high viral load (>10⁴ copies/ml) | 22          | 16 (72.7%)  | 13 (59.1%)  | 19 (86.4%)         |
| Single HBoV infection            | 36            | 26 (72.2%)  | 19 (52.8%)  | 33 (91.7%)         |
| Multiple viral infections        | 43            | 19 (44.2%)  | 21 (48.8%)  | 30 (69.8%)         |
| Total HBoV-positive              | 79            | 44 (55.7%)  | 40 (50.6%)  | 63 (79.7%)         |
mainland China and Hong Kong in children with LRTI.28,29 The limited number of specimens studied, however, does not preclude the possible presence of the ST1 genotype of HBoV. Interestingly, higher HBoV positivity and load were observed in throat swabs (n = 96) than in nasal swabs (n = 78), suggesting an infection preferentially of the lower respiratory tract, as indicated by previous studies on bronchoalveolar lavage (Fig. 4).11,30,31 In addition, HBoV was detected in sera of infected children, as previously observed with other parvoviruses.1 Consistent with other reports, samples with high copies of HBoV detected in nasal and throat swabs were more frequently positive in sera, compared with those with low copies5,6,20,24 suggesting that viremia often occurs during acute infection.

A high multi-infection rate was also detected in HBoV-infected patients (51.0%), with RSV and HRV having the highest prevalence. The high ratio of these two co-infecting viruses may reflect the possible presence of the ST1 genotype of HBoV. Interestingly, higher HBoV positivity and load were observed in throat swabs (data not shown). HBoV infection may be maintained to low copies in some individuals by an as yet unresolved interfering process triggered by the presence of the co-infecting agent. Whether HBoV exacerbates other viral infections or appears in children infected with other viruses is difficult to address and would require further studies.

Previous studies reported that the majority of epitopes recognized by antibodies in HBoV-infected patients were located in the VP2 protein. In this study, we divided VP2 into 4 fragments (P1–P4) and tested their antigenic reactivity by Western blot. Sixteen of 20 sera contained IgG/IgM antibodies that reacted with VP2 and its P3 fragment, and four did not show reactivity with VP2 and any of the four fragments. Strong immunoreactivity with IgG and IgM was observed for fragment P3, consistent with results with the whole VP2 protein. However, preparation of large amounts of purified P3 antigen in E. coli is easier and cheaper than that of VP2, which requires a more tedious preparation in the baculovirus system. Therefore, the purified P3 fragment of VP2 represents an alternative antigen for IgG detection in ELISA to test for acute or recent infection. It remains to be determined, however, whether this polypeptide is still recognized by long lasting antibodies to prevent false negative result for past HBoV infection.20,24,33 The number of IgG-positive cases increased with the age of the children, suggesting that infection of HBoV likely occurred during childhood. The data validate other studies that showed a high number of IgG-positive serum samples in adults. Since some previous studies reported the absence or only few cases of anti-HBoV IgM in virus-negative samples from children, and rare positive results in healthy adults,24,25 only sera of patients confirmed with HBoV infection were tested by Western blot for IgM. Anti-HBoV IgM antibodies were likely detected more often in patients viremic for HBoV or with high copies of HBoV in nasal and throat swabs, as previously observed. In 11 patients with viremia, high copy number in swabs, and infected only by HBoV, 10 (90.9%) were IgM-positive, which is consistent with previous studies.20,25 Patients infected singly with HBoV had comparatively more anti-HBoV IgM and IgG than multi-infection ones (p = 0.0119) (Table 4). In contrast, in 57 patients with low viral load in swab samples, the amount of IgG or IgM remained at a low level (data not shown).

Persistent HBoV shedding was reported in a few patients with symptoms lasting for several weeks,9,25 and in immunocompromised children. In this study, 12 patients were called for a second visit within an interval of 8–31 days (Table 5). No fever or symptoms of the respiratory tract were noted in these patients. Low copies of HBoV were still found in nasal and throat swabs, serum, and urine, indicating that infection of HBoV may last for a long period of time at a low level, and in the presence of anti-HBoV antibodies (Table 5). This situation was also observed in parvovirus
Systemic and long-term persistent infection, B19 infection with systemic and long-term persistent infection, high IgG reactivity and activated CD8+ T cell responses. 36 All ten B19 infection with systemic and long-term persistent infection, NA: Not available.

Persistent HBoV shedding in patients, with the interval between two visits.

Table 5

| Patient ID | Age / gender | Days interval | Nose swab | Throat swab | Serum | Urine | Antibody | Co-infection |
|------------|--------------|---------------|-----------|-------------|-------|-------|-----------|--------------|
| R3071260003 | 17 M/F       | 14 days apart | 3.2 × 10^4 | 3.2 × 10^4 | 3.0 × 10^4 | NA    | –        | –           |
| R3071203001 | 12 M/M       | 14 days apart | 2.3 × 10^4 | 4.6 × 10^4 | 9.6 × 10^4 | NA    | +        | –           |
| R3080218002 | 10 M/M       | 14 days apart | 5.0 × 10^3 | 9.0 × 10^3 | 1.4 × 10^3 | 0      | 0        | ++ HRV      |
| R3080218005 | 5 y/M        | 15 days apart | 6.1 × 10^2 | 2.9 × 10^2 | 0      | NA    | +        | ++ IAV      |
| R3080303002 | 13 M/M       | 8 days apart  | 0          | 2.1 × 10^2 | 5.3 × 10^2 | 0      | +        | ++ AdV      |
| R3080303004 | 5 y/M        | 20 days apart | 2.2 × 10^2 | 2.9 × 10^2 | 0      | NA    | –        | –           |
| R3080320004 | 5 y/M        | 31 days apart | 1.5 × 10^3 | 2.4 × 10^4 | 7.9 × 10^2 | 0      | 2.0 × 10^3 | +           |
| R3080508002 | 7 m/M        | 26 days apart | 1.6 × 10^2 | 3.4 × 10^2 | 0      | NA    | –        | + HRV       |
| R3080508005 | 9 m/M        | 26 days apart | 2.9 × 10^2 | 1.9 × 10^2 | 0      | NA    | –        | – PIV4/HRV  |
| R3080522002 | 5 y/M        | 12 days apart | 0          | 7.5 × 10^2 | 0      | NA    | –        | –           |
| R3080811001 | 11 m/M       | 20 days apart | 2.5 × 10^2 | 2.4 × 10^5 | 5.6 × 10^4 | 0      | 0        | + ++        |
| R3080825005 | 2 y/M        | 17 days apart | 5.0 × 10^5 | 4.5 × 10^5 | 7.5 × 10^4 | 0      | 1.2 × 10^3 | 0 ++        |

NA: Not available.

* m, Month; y, year.
** F, Female; M, male.
† HBoV load was normalized by GAPDH.

B19 infection with systemic and long-term persistent infection, high IgG reactivity and activated CD8+ T cell responses. 36 All ten patients with persistent PCR positivity of HBoV were found to be free of respiratory disease, but three were diagnosed with a secondary infection with ADV or HRV, suggesting that the persistence of HBoV was of less clinical significance, but may have played a role in promoting co-infection. Altogether, our study suggests that acute infection of HBoV causes systemic infection, induces immune responses, and may play a crucial role in respiratory disease of children, and is often associated with co-infection. Further studies are needed to understand the mechanism of the interactions between the host immune response, HBoV infection and prolonged virus shedding.

Conflict of interest

The authors declare no conflict of interest.

Acknowledgments

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