# PHYLOGENETIC ANALYSIS OF HUMAN BOCAVIRUS IN CHILDREN WITH ACUTE RESPIRATORY INFECTIONS IN IRAN

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Human bocavirus (HBoV) was first characterized in nasopharyngeal aspirates from young children with acute respiratory infections. It is prevalent among children with acute wheezing. This study was carried out in order to analyze the infection frequency and coinfection rates of HBoV with respiratory syncytial virus (RSV) and to perform phylogenetic analysis of HBoV in samples of children with acute respiratory infection in Isfahan, Iran. During the time period 2016–2017, altogether 75 respiratory samples from children hospitalized with acute respiratory infection were collected. The samples were first screened for RSV by direct immunofluorescence method and then subjected to detect HBoV DNA by PCR. Genotyping of HBoV-positive samples was conducted by direct sequencing of PCR products using NP and VP1/VP2 genes. Out of 75 respiratory samples, 20 (26.7%) and 10 (13.3%) were positive for RSV and HBoV, respectively. The coinfection rate was 40% (p = 0.048). Considering the seasonal distribution, winter has the highest extent outbreak (p = 0.036). Sequence analysis of positive samples exhibits that all of the isolated HBoV were related to genotype 1 (HBoV-1) with minimal sequence variations. Increasing frequency of HBoV suggests that the virus is related to acute respiratory infection in children. A single genetic lineage of HBoV1 seems to be the major genotype in Iran.

Keywords: human bocavirus, acute respiratory tract infection, respiratory syncytial virus, phylogenetic analysis, Iran

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## Introduction

Acute respiratory tract infections (ARTIs) are the frequently diagnosed diseases for hospitalization, particularly in infants and young children [1, 2]. The disease is the main cause of morbidity and mortality, estimating 3.5 million deaths worldwide [2]. Viral pathogens play an important role in ARTI [3]. A heterogeneous group of respiratory viruses, including some of the most notable RNA viruses, such as respiratory syncytial virus (RSV), influenza virus, parainfluenza virus, metapneumovirus, rhinovirus (RhV), and coronavirus are responsible for acute respiratory illnesses. RSV is the most prevalent virus involved in ARTIs [1-3]. Human bocavirus (HBoV), a member of Parvoviridae family, was first identified in 2005 in Sweden using a protocol based on DNase treatment, random polymerase chain reaction (PCR) amplification, intensewidely sequencing, and bioinformatics interpretation [4]. Its classification as bocavirus is based on similarities of the genetic structures and amino acid sequences with bovine parvovirus and canine minute virus [5-7]. PCR techniques enable detection of HBoV genome fragments from nasopharyngeal aspirates, bronchoalveolar lavages, feces, serum and urine samples using amplification of NP1, NS1 or/and the VP1/2 gene, or other nucleic acid-based determination methods [8-10]. HBoV clinical manifestations might vary in infants and toddlers and include fever, wheezing, coughing, and rhinorrhea [11, 12]. There are four major HBoV genotypes, such as HBoV 1-4 [13]. HBoV-1 has been mainly identified in respiratory samples, whereas HBoV 2-4 are chiefly identified in stool samples, indicating to cause gastrointestinal infections [13]. HBoV has a worldwide distribution [7, 13]. The virus is extensively circulating throughout the year, although it is more predominant in spring and winter [7, 13, 14]. The average prevalence of HBoV is 1.0%-56.8% in respiratory samples and 1.3%-63% in stool specimens of children with acute gastroenteritis [5, 13]. Coinfection between HBoV and other respiratory pathogens, such as RSV, and human RhV is frequent. Infants with coinfections (RSV and HBoV) have shown higher clinical severity than infants with RSV, RV, and BoV infections [15-17]. Notably, coinfecting pathogens have been found in up to 83% of respiratory samples [18]. The most typical age for HBoV infection is less than 2 years; rarely, it has been found in adults and the older people [7, 13]. The main goals of this report were to find the frequency of HBoV in children less than 3 years, to characterize its seasonal distribution, and to carry out phylogenetic analysis of HBoV strains circulating in Isfahan, Iran. We also studied the coinfection rates of the virus with the most prevalent respiratory pathogen, RSV.

### **Materials and Methods**

#### Patients and clinical samples

This cross-sectional study was carried out between 2016 and 2017 at teaching hospital, which is affiliated to Isfahan University of Medical Sciences (Imam Hussein Central Pediatric Hospital). Nasopharyngeal swabs collected from children less than 3 years old with symptoms of respiratory tract infection, including fever, wheezing, coughing, hypoxia, dyspnea, and rhinorrhea. The specimens were immediately transported in cold boxes (2–8 °C) to the virology laboratory. The nasopharyngeal swabs then divided into two aliquots: one treated for nucleic acid extraction and the second was immediately centrifuged at 3,000 rpm for 10 min to pellet the infected cells and stored at -70 °C [19].

#### Detection of RSV

For detection of RSV antigens in respiratory samples, direct immunofluorescence technique [20] was performed using RSV commercial kit (Euroimmun RSV, USA), according to the manufacturer's instruction. Briefly, secretions from the nasopharyngeal region were collected into a mucus extractor through a size 8 feeding tube. The mucus extractor and tubing were transmitted to the laboratory for processing. In order to dilute the mucus, 2 ml phosphate buffered saline (PBS) was added to the tube and centrifuged for 10 min at  $380 \times g$ . The pellet was suspended in 0.5 ml PBS by gently pipetting and 25 µl of the resuspended cell deposit was added to the slide. The specimen was air-dried thoroughly and fixed with fresh acetone at room temperature for 10 min. The fixed slides were stained with conjugated purified murine monoclonal antibodies directed against the fusion protein and nucleoprotein of RSV, mounted, and visualized through epifluorescence microscope (Olympus EX51, Japan).

# Detection of HBoV

Viral nucleic acids were extracted from respiratory secretion using viral nucleic acid extraction kit (DynaBio<sup>TM</sup>, Iran), according to manufacturer's protocol. All samples were screened for HBoV using PCR method by primers detecting the NP-1 coding region. The primers sequences were as follows: F: 5'-AGCTCTGTAAGTACTATTAC-3' and R: 5'-CTCTGTGTTGACTGAATA-CAG-3' [21]. PCR condition was as follows: initial denaturation at 94 °C for 5 min, followed by 40 cycles of denaturation at 94 °C for 1 min, annealing at 54 °C MOHAMMADI ET AL.

for 1 min, extension at 72 °C for 2 min, and a final extension at 72 °C for 10 min. The amplicons were visualized on 1.5% agarose gel stained with SYBR Safe DNA gel stain (ThermoFisher Scientific, USA). A cloned plasmid pTZ, containing HBoV NP-1 gene (Invitrogen, Carlsbad, CA, USA), was confirmed by sequencing and used as positive control.

#### HBoV genotyping

For genotyping, DNA of all HBoV positive samples was amplified by conventional PCR assay. Briefly, a set of forward (5'-GGACCACAGTCATCA-GAC-3') and reverse primers (5'-CCACTACCATCGGGCTG-3') were used to target 820 bp of the HBoV VP2 gene junction [22]. The thermal cycling conditions were as follows: an initial denaturation of 5 min at 94 °C, followed by 40 cycles of 1 min at 94 °C, 1 min at 54 °C, 2 min at 72 °C, and the final extension of 10 min at 72 °C. The PCR products were purified and subjected for sequencing, (Bioneer Inc., Korea).

## Phylogenetic analysis

All alignments were performed using Chromas Pro software program (version 2.1.3; Techelysium Pty Ltd., South Brisbane, QLD, Australia). Phylogenetic trees of the NP-1 and VP-2 genes generated using the neighbor-joining and maximum likelihood methods and Kimura 2-parameter algorithm of MEGA software (version 6.0; Pennsylvania, USA). Bootstrap probabilities for 1,000 literations were calculated to check confidence estimates. Only those samples that showed the same genotype in both methods were included for inference. The nucleotide sequences of NP1 and VP1/VP2 sequences were compared with those of reference strains available in NCBI website using BLASTN. The partial sequences of VP1/VP2 gene were submitted to GenBank (accession numbers: MH574941–MH574945).

#### Statistical analysis

Data were processed using SPSS statistical software program version 16.0 (IBM, SPSS statistics, Chicago, IL, USA). The correlations were subjected to the Pearson's  $\chi^2$  test. Statistical significance is set at a *p* value of less than 0.05.

## Ethical approval

This work was in accordance with the ethical standards of Ethics Committee of the Isfahan University of Medical Sciences (9604-396486). Informed parental consent was obtained from all participants.

#### Results

#### Detection of RSV

In total, 75 respiratory samples were collected from children less than 3 years old admitted to the referred hospital in Isfahan, Iran. Twenty (26.7%) of the specimens were positive for RSV (Figure 1). Twelve (60%) positive samples were collected from females and 8 (40%) were collected from males. No statistical significance was found regarding RSV infections and gender (p = 0.512). The age distribution showed that 7 (35%) of RSV-positive samples were from children between 1 and 2 years old. The statistical significance was meaningful regarding RSV and age (p = 0.032). The prevalence of ARTIs caused by RSV has a major peak in winter, especially in January and February (Figure 2). In this study, 11 positive cases (55%) were collected in winter. There was meaningful statistical significance regarding the RSV infection and season (p = 0.039).

#### Detection of HBoV

Presence of HBoV in specimens was evaluated by PCR method. Among the 75 collected respiratory samples, HBoV DNA was found in 10 (13.3%) samples. The size of the PCR product including the fragment of the NP1 gene was 354 bp (Figure 3). There was no statistical significance regarding HBoV infection and gender (p = 0.506). Among HBoV-infected children, 40% were between 1 and 2 years old, followed by 30% under 6 months, 0%, between 6 months and 1 year,



Figure 1. Epithelial cells in nasopharyngeal swabs specimen from a positive control (A) and patient with RSV infection (B), showing specific RSV fluorescence after monoclonal antibody staining against F/NP. White arrows show the staining of prominent RSV inclusions (40×)

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Figure 2. Monthly distribution of HBoV, RSV, and RSV/HBoV coinfection. HBoV: human bocavirus; RSV: respiratory syncytial virus (p < 0.025)



**Figure 3.** Analysis of NP-1 PCR products by agarose gel electrophoresis. Viral nucleic acids extracted from respiratory secretion were amplified using PCR. Lanes 2 and 7: two positive samples; Lanes 1, 4–6, and 8: five negative samples; negative control and positive control. NP-1: nucleoprotein 1

and 30% between 2 and 3 years (p = 0.037). Among 10 HBoV-positive samples, 9 (90%) were collected in winter. This showed that the prevalence of HBoV infections has a major peak in winter (p = 0.036; Figure 2).

# RSV and HBoV dual infections

Comparing the patient groups according to the type of virus (RSV, HBoV, and HBoV + RSV coinfection), several differences were identified between the groups. Mean age of children was different between RSV and HBoV mono-infection group and RSV/HBoV coinfection. Children with RSV and HBoV single infection were older than RSV/HBoV coinfection group (p = 0.037). Clinical symptoms such as fever and rhinorrhea were not significant (p > 0.05). There was meaningful statistical significance regarding clinical symptoms, including wheezing (p = 0.002), cough (p = 0.021), dyspnea (p = 0.048), and HBoV infection (p = 0.048). Dyspnea (40%; p = 0.002) was the most common symptom in the HBoV group and coughing (90%; p = 0.021) was the most frequent symptom in RSV/ HBoV coinfection group (Table I).

## Phylogenetic analysis

Two PCR products of NP-1 gene were confirmed by sequencing. The results of sequences were compared with 34 reference sequences including strains from

Clinical features	RSV $(n = 75)$	HBoV $(n = 10)$	RSV + HBoV (n = 4)	p value
Male	8 (40%)	5 (50%)	1 (25%)	0.506
Female	12 (60%)	5 (50%)	3 (75%)	
37.9 < fever	19 (95%)	5 (50%)	3 (75%)	0.469
Wheezing	10 (50%)	7 (70%)	3 (75%)	0.002*
Cough	16 (80%)	9 (90%)	3 (75%)	0.021*
Rhinorrhea	15 (75%)	7 (70%)	4 (100%)	0.407
Hypoxia (SatO <sub>2</sub> < 95%)	5 (25%)	4 (40%)	2 (50%)	0.105
Dyspnea	4 (20%)	4 (40%)	1 (25%)	0.048*
Age: <6 months	5 (25%)	3 (30%)	2 (50%)	0.037*
Age: 6 months-1 year	2 (10%)	0	0	
Age: 1-2 years	7 (35%)	4 (40%)	1 (25%)	
Age: 2–3 years	6 (30%)	3 (30%)	1 (25%)	

 Table I. Clinical and demographic characteristics in children with acute respiratory infections regarding the identified viruses

Note: RSV: respiratory syncytial virus; HBoV: human bocavirus.

\*p values <0.05 were considered to be statistical significant.



Figure 4. Phylogenetic tree based on gene sequences encoding the HBoV polyprotein NP1 of HBoV isolates. The red triangle represents sequences originating from our studies (MH377116 and MH061315). HBoV1-4 standard genotypes were represented in squares: Black is related to HBoV-1, purple related to HBoV-2, turquoise related to HBoV-3, and blue related to HBoV-4 genotypes. The percentage of replicate trees in which the taxa are clustered in test (1,000 replicates bootstrapping) is shown next to the branch

the four HBoV genotypes registered in the GenBank database (https://www. ncbi.nlm.nih.gov/genbank/). Phylogenetic analysis of the partial NP1 gene confirmed that these two isolates are HBoV and belong to genotype 1 (Figure 4). The sequenced NP-1 gene was submitted to GenBank with following accession numbers: MH377116 and MH061315. Because VP1/VP2 gene has the greatest variation in the HBoV genome, especially at its 3' end [23], all HBoV-positive specimens were selected to amplify VP1/VP2 partial gene sequences and aligned with 44 reference sequences including HBoV1-4, porcine, bovine, and canine parvoviruses. Phylogenetic analysis of VP1/VP2 region indicated that all HBoV isolates in this study in the isolated strains were in the same cluster as other strain from Iran (Figure 5).



Figure 5. Phylogenetic tree constructed from the VP1/VP2 gene sequences of HBoV strains from respiratory samples. The red triangles represent strains from children with acute respiratory infection (GenBank accession numbers: MH574941, MH574942, MH574943, MH574944, and MH574945)

#### Discussion

Since the discovery of HBoV by Allander, rising reports have shown that HBoV is involved in upper and lower respiratory tract infections [4, 6]. HBoV has been detected in 1.5%–19% of respiratory tract samples from young children with respiratory infections [13]. Reports from New Zealand, North America, Europe, Asia, and Africa confirm the global distribution and indicate increasing frequency of the virus worldwide [24–27]. In this study, HBoV was detected in 10 of 75 (13.3%) respiratory samples. In accordance with this study, the prevalence of HBoV infections in some regions of Iran was also evaluated. The prevalence has been reported 8% in a northern city of Iran (2003–2004) [28], 6.8% in Tehran (2006–2008) [29], and 10.7% (2012–2013) in Tehran [30]. The minor difference in frequency could be related to the number of patients studied, time of collecting the samples, geographical locations, and the age groups studied. In addition, it seems that the prevalence and distribution of the virus is increasing. Although

RSV is the major pathogen in infants less than 1 year old, the rate of HBoV infection was also meaningful and the pathogenic potential of this virus could not be ignored.

The most common symptoms of acute viral respiratory infections are wheezing, cough, dyspnea, and fever [31]. HBoV is prevalent among children with acute wheezing [8, 12, 13]. We reported that major symptoms in RSV, HBoV mono-, and coinfections were different. The most frequent clinical manifestations in HBoV group were wheezing and dyspnea. This could explain why pneumonia is much more frequent in children infected with HBoV [12–14]. RSV and HBoV coinfection shared clinical symptoms of both viruses, with wheezing as the most frequent clinical symptom. This result is in agreement with other studies that outline HBoV-1 is able to infect the lower respiratory tract down to bronchioles [12].

The acute respiratory infections usually occur from late fall to early spring [7, 13]. Seasonal peaks of HBoV infection vary among different countries because of different climate and geographic conditions. Previous studies suggested that HBoV infection had a higher detection rate in winter [7, 12, 13]. In this study, a higher frequency of HBoV infection was observed between January and March with a peak in February. Bocavirus is often found in mixed infections with other viruses [15]. Prevalence of HBoV seasonally coincides with RSV [15–20]. Therefore, we studied RSV as one of the most commonly associated infectious agents in ARTI. We reported the coinfection rates with RSV [28, 29], whereas in other countries, a high proportion of coinfections with other viruses have been observed [27, 31, 32]. This considerable coinfection rate further supported the idea that HBoV might be a bystander in respiratory infection.

Based on Vp1/VP2 gene sequences, all HBoV strains isolated in this study were related to HBoV-1. Although very low variability exists in HBoV genomic region encoding capsid proteins, all our isolates were in the same cluster as the original isolates of ST1 (Stockholm 1; 5,217 nt; accession no. DQ000495) and ST2 (Stockholm 2; 5,299 nt; accession no. DQ000496) [4]. Moreover, other HBoV strains identified in Iran and some countries in the region are in the same cluster (Figure 5) [30, 32]. This proposes that relative HBoV-1 strains are circulating in this geographical region.

According to our results, the frequency of HBoV infections/circulation in Iran is increasing. This virus might play a significant role in ARTIs, as a major cause or coinfection with RSV. All the strains detected in respiratory tract infections belonged to HBoV genotype 1. Further studies on the clinical relevance of HBoV in respiratory tract infections are needed.

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#### **Conflict of Interest**

The authors have no potential conflict of interest to declare.

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