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Complete genome sequence of human astrovirus genotype 6

Li Guo¹, Richard Gonzalez^{1,2}, Wei Wang¹, Yongjun Li^{1,2}, Gláucia Paranhos-Baccalà², Guy Vernet², Jianwei Wang^{1*}

Abstract

Background: Human astroviruses (HAstVs) are one of the important causes of acute gastroenteritis in children. Currently, eight HAstV genotypes have been identified and all but two (HAstV-6 and HAstV-7) have been fully sequenced. We here sequenced and analyzed the complete genome of a HAstV-6 strain (192-BJ07), which was identified in Beijing, China.

Results: The genome of 192-BJ07 consists of 6745 nucleotides. The 192-BJ07 strain displays a 77.2-78.0% nucleotide sequence identity with other HAstV genotypes and exhibits amino acid sequence identities of 86.5-87.4%, 94.2-95.1%, and 65.5-74.8% in the ORF1a, ORF1b, and ORF2 regions, respectively. Homological analysis of ORF2 shows that 192-BJ07 is 96.3% identical to the documented HAstV-6 strain. Further, phylogenetic analysis indicates that different genomic regions are likely undergoing different evolutionary and selective pressures. No recombination event was observed in HAstV-6 in this study.

Conclusion: The completely sequenced and characterized genome of HAstV-6 (192-BJ07) provides further insight into the genetics of astroviruses and aids in the surveillance and control of HAstV gastroenteritis.

Background

Human astroviruses (HAstVs) are one of the most common causes of acute gastroenteritis in children worldwide [1-3]. HAstV was first identified during an outbreak of gastroenteritis among hospitalized infants in 1975 [4]. Its name is derived from its distinctive star-shaped appearance under the electron microscopy (EM). Molecular analyses indicate that HAstVs are non-enveloped viruses with a 6-8 kb single-stranded, positive-sense RNA genome consisting of three overlapping open reading frames (ORFs)—ORF1a, ORF1b and ORF2—as well as the 5'- and 3' nontranslated regions (NTRs) [5]. ORF 1a encodes a serine protease; ORF 1b encodes an RNA dependent polymerase; and ORF 2 encodes a capsid precursor protein [5].

HAstVs have been grouped into eight known serotypes (HAstV-1 through HAstV-8) based on their reactivity to polyclonal antibodies and on analysis by immunofluorescence assays, neutralization assays, and

immunolectron microscopy (IEM) [5-7]. Phylogenetic analyses of the HAstV nucleotide sequence have defined eight genotypes, and further studies have indicated a strong correlation between the genotypes and serotypes [8]. As such, genotypes are frequently applied to type HAstVs.

Genomic characterization studies are important to the understanding of the origin, molecular evolution, and phylogenetic relationships among HAstV genotypes. The full-length genome sequence for a HAstV (HAstV-2) was first determined in 1993 [9]. Subsequently, the complete genomic sequences of five more genotypes (HAstV-1, HAstV-3, HAstV-4, HAstV-5, and HAstV-8) were reported [9-12]. Because the dominant, disease-causing HAstV type and strain often fluctuate with time and geographic location, it is critical that we characterize the complete genomic sequences of all known genotypes in order to better control and prevent future epidemics [13]. Limited sequence information for HAstV genotype 6 is available. Only a partial genome sequence has been reported [14,15], even though this genotype has been identified as one cause of sporadic or large scale outbreaks of acute gastroenteritis worldwide [16,17].

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In 2007, we identified a case of HAstV-6 infection in Beijing, China, suggesting that this strain might be more epidemiologically relevant than previously recognized [18]. Here we sequenced and analyzed the complete genomic sequence of this HAstV-6 192-BJ07 strain, and describe its genetic characteristics by comparing its sequence with other known HAstV genotypes. The characterization of HAstV-6 by whole genome sequencing provides critical insight into the genetics of this virus as well as valuable information for the control and prevention of HAstV-induced gastroenteritis.

Results

Genome organization

Complete genome sequencing of HAstV-6 (192-BJ07) was performed from a stool sample that tested positive for HAstV-6 RNA fragments by RT-PCR. Starting with the cloning of ORF2, the entire sequence of the viral genome was obtained by a step-wise amplification strategy through 5'- and 3'-RACE. The full-length genomic RNA of 192-BJ07 is 6745 nucleotides (nt) in length, excluding a poly-A tail at the 3' end. HAstV-6 (192-BJ07) has the same genome organization as other known HAstV genotypes. It has a 5'-NTR of 82 nt, a 3'-NTR of 81 nt, and three overlapping major ORFs: ORF1a (2766 nt), ORF1b (1548 nt), and ORF2 (2337 nt). Details of the predicted genome organization of 192-BJ07 are shown in Fig. 1.

ORF analysis

The sequence of the HAstV-6 192-BJ07 strain displayed similarity to those of other known HAstV genotypes. ORF1a of the HAstV-6 192-BJ07 strain shared 79.0%

79.9% nucleotide identity and 86.5-87.4% amino acid identity with those of genotypes 1 through 5 and with genotype 8. Two mutation sites were found at amino acids 757 and 758 in 192-BJ07 ORF1a, which result in the insertion of Arg and Lys.

Unlike ORF 1a, ORF1b exhibited higher nucleotide (85.7%-87.5%) and amino acid (94.2-95.1%) identity with that of genotypes 1 through 5 and genotype 8 (Table 1), indicating that ORF1b is more conserved among the 192-BJ07 strain and the HAstV genotypes than ORF 1a.

Pairwise comparisons of the nucleotide and amino acid sequences of the ORF2 region showed that 192-BJ07 shares relatively low identity with other known HAstV genotypes in this region (62.4-72.6% nucleotide identity, and 65.5-74.8% amino acid identity; Table 1). However, 192-BJ07 exhibited high identity—96.3% nucleotide identity and 95.9% amino acid identity—with the documented sequence of the HAstV-6 strain (GenBank accession number Z46658 and Table 1). Structural predictions of ORF2 indicated that there are three highly conserved amino acid residues that can be cleaved to yield proteins with different sizes: Lys 71 for a 79-kDa protein, Arg 361 for VP29 and Arg 395 for VP26 [19].

Alignment of the 192-BJ07 ORF2 sequence with the seven other HAstV genotypes showed that the conserved domain (N-terminal 415 amino acids) shares 85.8-89.6% identity (Fig. 2); whereas the region from amino acid 416 to the carboxyl-terminus, i.e. the variable region, showed higher levels of variation, with amino acid sequence identities of only 41-58% (Fig. 2). Compared with other HAstV genotypes, the 192-BJ07

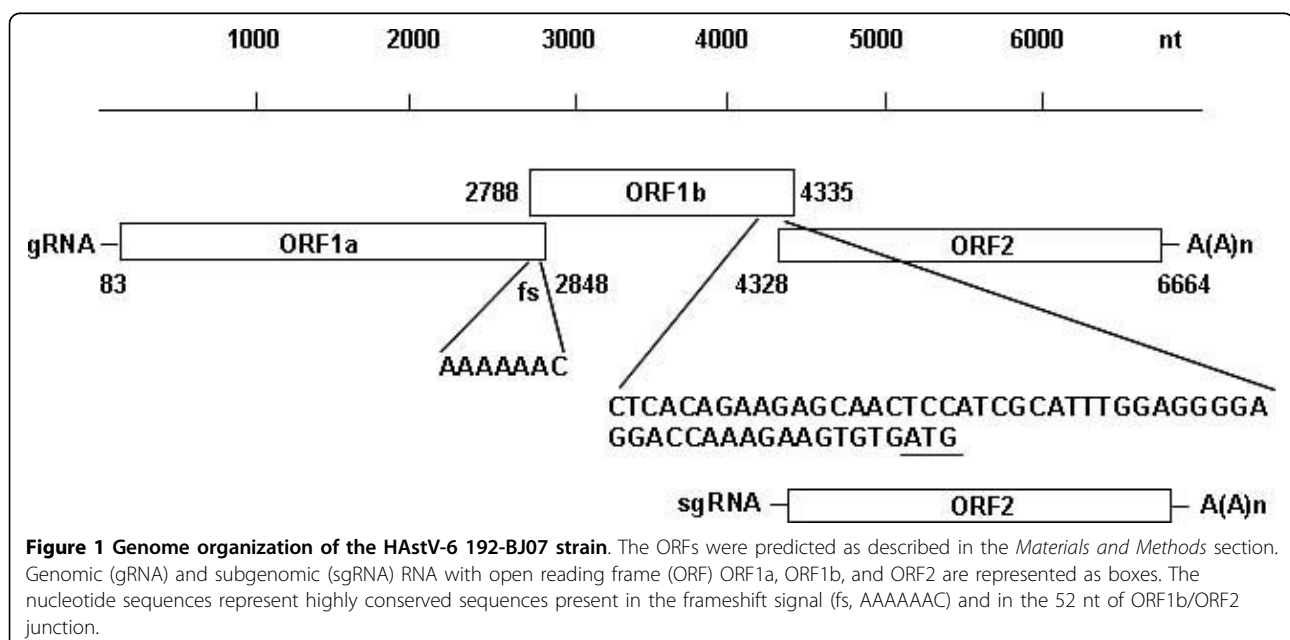


Table 1 Sequence identity between HAstV-6 (192-BJ07) and other HAstV genotypes

Genotype (Genbank accession numbers)	Percent identity							
	5'NTR	ORF 1a		ORF 1b		ORF 2		3'NTR
	Nucleotide	Nucleotide	Amino acid	Nucleotide	Amino acid	Nucleotide	Amino acid	Nucleotide
HAstV-1 (L23513)	76.5	79.3	87.1	86.4	94.8	71.3	71.4	98.8
HAstV-2 (L13745)	42.7	79.4	87.3	86.0	94.2	69.9	67.5	95.1
HAstV-3 (AF141381)	58.5	79.0	86.6	85.7	94.6	72.6	72.8	96.3
HAstV-4 (AY720891)	76.8	79.9	87.0	86.5	95.1	62.4	65.6	92.6
HAstV-5 (DQ028633)	57.3	79.2	86.5	87.5	94.6	72.1	74.8	96.3
HAstV-6 (Z46658)	NA	NA	NA	NA	NA	96.3	95.9	NA
HAstV-7 (Y08632)	NA	NA	NA	NA	NA	72.4	72.3	NA
HAstV-8 (AF260508)	57.8	79.8	87.4	86.0	94.8	70.9	72.8	97.5

NA: Not available.

strain displayed 25-57.5% variation in VR1 (aa 292-319) and 30.8-92.3% in VR2 (aa 386-399) [20] (Fig. 2). These results are consistent with those described in a previous report on HAstV-8 by Méndez-Toss et al [20,21].

Non-coding region analysis

HAstV genotypes typically contain 80 to 85 nt in the 5'-NTR. The region shows the highest nucleotide variation (23.2-57.3%) among the five regions of the viral genome (5'-NTR, ORF1a, ORF1b, ORF2, and 3'-NTR) (Table 1). However, the 37 nt of the very 5' end of the viral genome are highly conserved (Fig. 3). Secondary structural predictions indicate that the HAstV-6 192-BJ07 strain has three stem-loop structures in the 5'-NTR. However, because there is a high level sequence variability within the region of nt 38-85 (Fig. 3), the consensus of the stem-loop structures is very low among HAstV genotypes (data not shown).

In contrast, we found that the nucleotide identities of the 3'-NTR sequences are as high as 92.6-98.8% compared with other known HAstV genotypes. However, the sequence variability within this region also results in secondary structure disparities of the 3'-NTR between HAstV genotypes (data not shown).

It has been recognized that HAstV RNA has a cis-acting element [ribosomal frameshifting heptamer sequence (AAAAAAC)] followed by a stem-loop structure in the ORF1a/1b junction region [22]. The 192-BJ07 strain also has such a shifty heptamer sequence and a similar stem-loop structure based on analysis with RNAstructure 4.5 software. This conservation may reflect the importance of such structures for translational regulation [22].

The ORF1b/ORF2 junction has been regarded as a regulatory element of the sub-genomic RNA (sgRNA) [23]. The alignment analysis of 52 nt at the ORF1b/ORF2 junction revealed that 192-BJ07 has a very high identity (98.4-100%) with other HAstV genotypes, consistent with a previous report [24].

Phylogenetic analysis

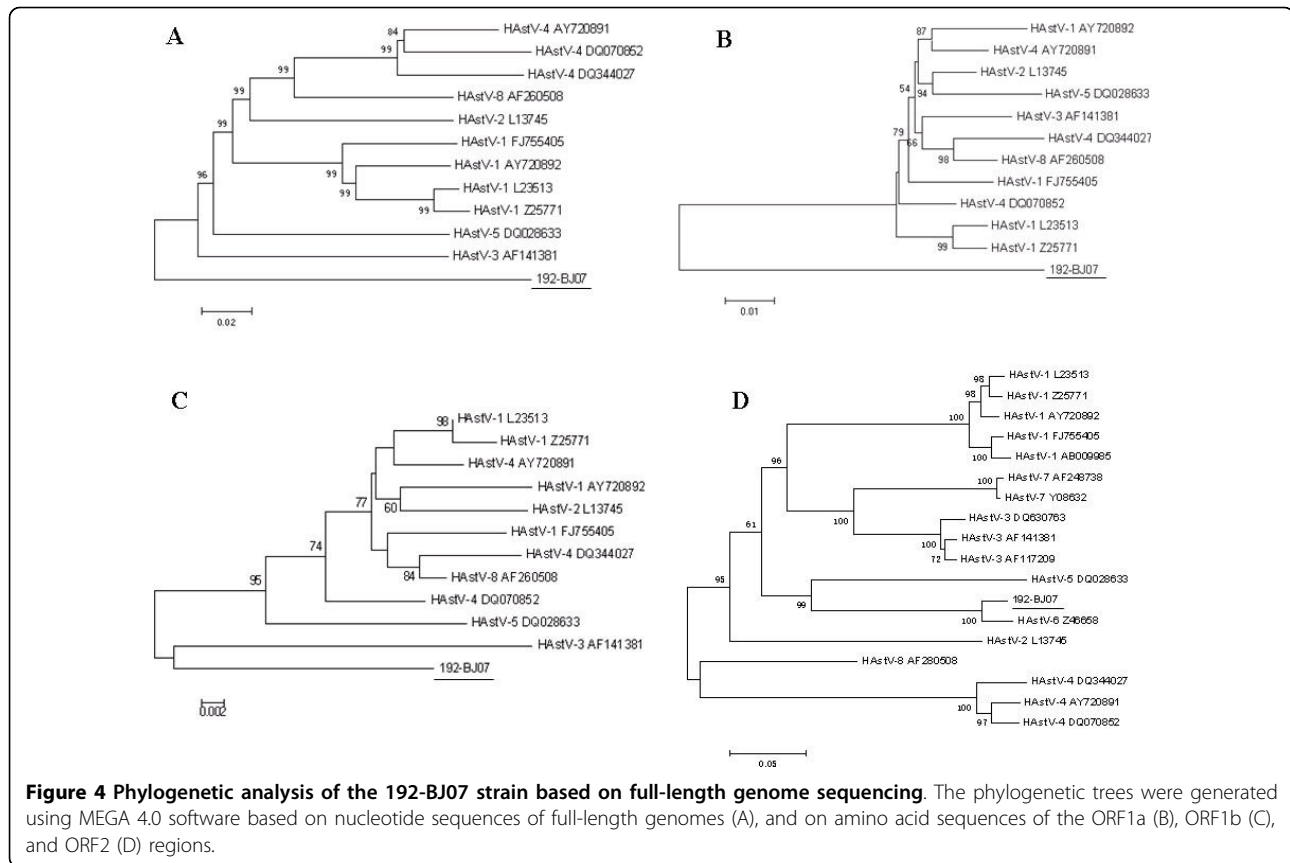
Nucleotide alignment analysis of whole genome sequences showed that the identities between the HAstV-6 192-BJ07 strain and the corresponding sequences of HAstV-1, -2, -3, -4, -5 and -8 were 77.2-78.0%. We found that the evolutionary relationships among HAstV genotypes were divergent when phylogenetic analyses were performed at the level of complete genome sequence or at the level of individual proteins. The phylogenetic analysis of the whole genome sequence and the ORF1a region indicates that the HAstV-6 192-BJ07 strain is a significant outlier on the phylogenetic tree compared to the other HAstV genotypes (Fig. 4A and 4B). In the region of ORF1b, the HAstV-6 192-BJ07 strain and HAstV-3 branch out earlier (Fig. 4C); while in the region of ORF2, HAstV-8 and HAstV-4 share the common ancestor role of other HAstV genotypes (Fig. 4D).

Recombination analysis

To determine whether there were recombination events between the 192-BJ07 strain and other known HAstV genotypes, we analyzed the similarities at the genome level. A similarity plot comparing the nucleotide sequences of HAstV-6 192-BJ07 and HAstV-1, -2, -3, -4, -5, and -8 is shown in Fig. 5. The sequence identities of the 192-BJ07 ORF1b region with the other genotypes

192-BJ07	MASKEIDMIVTEVNNR--GGRSISKRSRQSRGRGSRVLTIVNSQKGRQKGRMKBQSNRVRNIVNKLRLDQVGTGFRPAICQKATATLGTIGSNTGG	98
HastV-1N.....S.....N.....DK.....R.RA.....P.D.....S.....K.....R.....V.....S.....	98
HastV-2N.....H.....Y.....K.....K.....R.....T.....	98
HastV-3K.NN.....N.R.....K.....RSR.....D.Y.....T.....K.....R.....S.....	100
HastV-4S.....N.....Y.....K.....K.....T.....T.....	98
HastV-5PS.....S.R.P.....DK.....R.....M.....K.....R.....V.....S.....	97
HastV-6T.....K.....R.....S.....	98
HastV-7N.S.R.G.....F.K.R.....L.....K.....R.....S.....	99
HastV-8S.....H.....Y.....K.....K.....T.....	98
192-BJ07	TTEITACILLNPLVKDAGTGTQGFVQALGAQYSMWKLYLWRLTSMVGSANVGTVVRVSLNFTSPSTSWFGLGARKGLDVTGKNAIFKLFPSD	198
HastV-1K.....L.....T.....	198
HastV-2	A.....K.....L.I.....M.....R.V.....R.....	198
HastV-3K.....V.....A.....	200
HastV-4	A.....D.....I.....	196
HastV-5K.....S.....D.....V.....A.....	197
HastV-6E.....	186
HastV-7	S.....K.....S.A.I.....V.R.A.....	199
HastV-8	A.....I.....V.....	186
192-BJ07	LOGPFDQWLTWVNDASDGLPVSVEIHLGRTMSYQHQFTGGFLVELASWCFCTGTAAPHLVRLMDEQVHVTFEGSAGTFLMNVFAASHFA	298
HastV-1I.....K.E.....N.P.....S.I.....SEG.....	298
HastV-2I.....K.K.E.....H.E.....K.I.....AEH.....	298
HastV-3M.....K.E.....N.I.K.E.....	300
HastV-4I.....Q.....T.....G.S.A.....S.D.....I.....EH.....	298
HastV-5I.....M.....N.....E.....S.....T.....V.....	297
HastV-6I.....K.....L.....M.....S.....DN.....I.K.DT.....	298
HastV-7I.....Q.....T.....S.A.....S.....I.....EH.....	299
HastV-8I.....Q.....T.....S.A.....S.....I.....EH.....	298
192-BJ07	RTVAQRSTLPTSMARAGENTSIVVQVLNAVSAELVTPFFFNVLVQSGVFFVLLGARGVSGRSFYVYASTQDALSNEFALCTQSGMRTFAV	398
HastV-1LA.....T.TL.....R.....T.....T.T.S.....P.....G.....MP.....	398
HastV-2MAE.H.SIS.TFS.....GDA.....T.....A.....I.....T.T.IHQ.....P.....GVT.GV.L.TIP.....	397
HastV-3AVA.SV.LT.Q.....T.....A.T.T.....P.....A.SG.....A.....	400
HastV-4MAVEN.S.S.TLS.....GESS.....T.....A.T.A.R.P.....GVTSTY.QSNP.....	398
HastV-5A.VA.....L.....R.....T.....T.N.T.....P.....I.....GL.SGL.AAS.....	397
HastV-6A.....T.....N.....	398
HastV-7AVA.S.....TL.....Q.....T.....I.....I.....S.A.M.....P.....V.G.V.A.I.....	399
HastV-8AVEN.S.S.LS.....GESS.....T.....A.T.A.R.L.....GV.A.A.QSNP.....	398
192-BJ07	MTLQFTQMNQFSLGHGETFATLGRSIFTSGETLKVLLIIGHPLSPMETHNQVWKNKIEPFG-----AVKIGRQDAQNYCTMNGFLITKVDWFTFEFQ	493
HastV-1	T.....A.AI.....AP.EY.V.F.S.M.AN.....PLDA.SGH---YN.AK.VD.L.Q.....S.AS.....Y.ID.....	495
HastV-2	V.....HT.I.SIVQDFSGE.R.....V.SIM.....SADR.V.L.LTA.TNSNDNL.AH.LG.LI.Q.MH.KT.E.Y.PD.....	497
HastV-3	R.....MT.....A.SP.DNPN.V.....Q.LA.TLT.N.L.TVS.QNH---H.AK.VNR.T.Q.....P.SS.....Y.TD.....	497
HastV-4	R.....R.A.....EP.DQF.V.M.V.ALWQ.RSDT.N.LP.VT.TGH---DAARV.WNT.L.IQ.L.DSLE.L.PHL.....	495
HastV-5	A.....NT.....A.A.DR.....I.V.Q.VT.....N.....G.NT.TE---E.....VNT.N.VV.....M.SSI.YD.M.....	492
HastV-6V.....R.....T.....L.....	493
HastV-7	P.....VT.....P.D.F.V.....Q.LA.TL.N.....TTA.QQQ---H.AK.TSN.T.Q.....P.SN.T.Y.D.....	496
HastV-8	R.....A.M.F.....EP.QFR.....V.F.MA.TA.S.N.....V.ENQ---YT.....I.LE.T.Q.....PVES.S.Y.AD.....	495
192-BJ07	PSEEPAPVQGLMVLNDHQBADVTAAGQYKPIITNDKQVTSVFLVRYRNGFGVTRHRSYFYRNSVNTDAVENIKRSATRHITVFNQSGWILLISTVL	593
HastV-1A.IK.....Q.VHS.....VK.PVTAQ.N.H.....L.....K.TY.....N.YL.....A.ATG.TT.VL.GD.YTAGES.T.G.....N.SI.....	595
HastV-2RD.T.IA.MS.MVHIT.....FMK.P.SI.NRM.I.....IKPLAD.K.QCM.....K.E.HDN.G.A.LLV.M.SPE.I.QV.E.....T.....	597
HastV-3A.T.I.P.....Q.VNS.....I.L.NQ.N.V.L.I.....K.TTS.N.YL.....SYGTGT.....L.V.D.TAQD.N.PV.....M.AI.....	596
HastV-4	E.Q.PILPE.G.YIGI.....L.FM.....V.H.N.H.S.I.IKPT.N.S.....YM.....F.E.QSDQN.A.L.Pq.WQ.N.QR.K.....V.N.AI.....	595
HastV-5L.V.V.MSE.L.MEGIN.....S.S.N.H.I.....Y.....K.N.....L.....P.EK.D.T.T.LM.....Pq.YT.....N.V.NN.....	592
HastV-6G.....E.....L.....Y.....	593
HastV-7P.IS.....Q.V.SR.....V.L.HFS.T.D.L.I.....K.TTS.N.L.....AAGTGT.....F.....G.SEQNIS.SE.....M.N.AT.....	595
HastV-8D.S.I.P.....YARVN.T.....GV.F.SSH.NRM.I.....TTS.....I.YT.....I.GAESGM.S.L.....Dq.Y.PLQ.T.K.....M.....	595
192-BJ07	HTGFPYSQWLNWQELQNDQATIIDQTMHLITPPVPSQIVFEMAITVPTQTRKCEETGLELAMG--ISEDEH-PISHVDEEETETEYTSDEDEIDE	689
HastV-1	VD.AMP.V.N.V.KWT.NM.K.LV.....M.L.E.T.MCY.L.SI.RS.A.HGVESDW---IEYLDAPD-SADQKEDI.BTDI.T.....D.....	690
HastV-2	KEWLP.E.V.DRV.KS.TP.YA.ALTYP.....D.....L.GH.TL.KISSPDPFGYRMSHQWQDCSDDDDUCE.VS.....T.D.E.....D.....	697
HastV-3	FNA.AFP.I.Q.V.L.T.L.M.....M.ANT.LL.....R.SISGS.SMISSPEEFTHEFGDWCDAID-ASDSRWLL.....D.D.E.....D.....	695
HastV-4	RN.....P.V.D.I.T.ESI.YA.VLA.F.N.QN.K.....VH.TM.S.ARSIGLEEDQ-----TDWQEPD.DLATS.ESDY.....S.....	682
HastV-5SLP.....V.....M.NEN.....M.K.M.T.....L.L.SL.AMAREFDQ.DQAVVA---HDS---GQALFSA.....DSDF.T.....	685
HastV-6I.....L.....Sg.G.....V.....D.Y.....	689
HastV-7	FNP.AFP.I.K.V.D.NTP.....V.....M.....MS.....GT.LL.....K.T.SG.NVSHFDH.NFSP---VMCDALD-AADVVELPT.DT.S.E.....D.....	682
HastV-8D.TS.....V.....T.NI.EV.F.M.Y.....AA.L.LH.VL.A.SEEP.YVDAPLF-----EPPPIE.....DSDF.T.....N.....	684
192-BJ07	VDFEFDLCT--SISEDD-IESNRVILLSTLNQMTVDRAIMTSRAFF----TFNHSRRELSNDLLGFSDCIATARSACNACNES-----CQLSGSR	774
HastV-1	A.....IID.....E.G.H.TD.....V.....MT.....R.AR.....LSDEIK.GYIM.....VSYSPFGH.....E.AKAVGETHFC.....	783
HastV-2	A.....HSPYS.EP.....S-D.N.....E.....R.K.....CAE.LK.SYIM.....ASGAFSS.W.N.D.ARWVGSNQLAK.....D.....	792
HastV-3	A.....HSTYG.EP.....-D.N.....E.....R.K.....SAG.TK.SYIM.....ASGLSPGN.....E.ARMINGNTPWV.....D.....	790
HastV-4	LE-----DE.D.EDSNTC.ELVIN.V.....ISRE.....Y.GMS.Y.NVWVSGGQ.TSQHIGETSSIDVAGAHY.CV.ERE-----GQSLNQ.....	787
HastV-5HLS---E.D.DV.N.....S.E.....R.N.....RAARL.SVY.....VSLGFGA.....EQARKVGDHDL.A.....	779
HastV-6N.....V.....N.....Y.P.....P.....A.....T.....	774
HastV-7	A.....HSTG.EP.....-D.N.....E.....K.....CAD.GK.SYIM.....ASGLSPGNV.....E.ARMINGNTPWV.....D.....	787
HastV-8HPS---E.D.DV.N.A.....L.....IS.E.....R.NG.....RAARV.SVY.....VSLGFGA.....EQARRAGDHDL.....	778
192-BJ07	GHAE	779
HastV-1	788
HastV-2	796
HastV-3	794
HastV-4	771
HastV-5	783
HastV-6	778
HastV-7	791
HastV-8	D..	

Figure 2 Alignment of the complete ORF2 amino acid sequence of 192-BJ07 with other HAstV genotypes. The alignment analysis was performed using the MegAlign programs in the DNASTar software package. Variable regions VR1 and VR2 are underlined. Dots indicate amino acid sequence identities, and a dash denotes a deletion. The single-letter amino acid code is used. GenBank accession numbers: HAstV-1: L23513; HAstV-2: L13745; HAstV-3: AF141381; HAstV-4: AY720891; HAstV-5: DQ028633; HAstV-6: Z46658; HAstV-7: Y08632; HAstV-8: AF260508.



the replication and/or transcription level among HAstV genotypes. The fact that the 5'-end of the 5'-NTR and the 3'-NTR and the 52 nt region at the ORF1b/ORF2 junction are highly conserved points to their critical role in the interaction with the viral replicative or transcriptional machinery. The variation in the 3'-end of 5'-NTR may influence the efficiency of viral genome replication or transcription, resulting in a difference in replication ability or virulence among different genotypes or strains [5].

The -1 ribosomal frameshifting is critical for the translation of the astrovirus genome [22]. The -1 ribosomal frameshifting requires two cis-acting signals: a shifty heptamer sequence (AAAAAAC) and a potential stem-loop structure [10,26]. This study showed that the HAstV-6 192-BJ07 strain also has such cis-acting elements, and further demonstrates the conservation of such elements among HAstV genotypes [5].

At present, the mechanism of HAstVs' variations is unclear. One study has indicated that recombination may be responsible for HAstVs' variation [24]. However, current studies have not broadly established the role of recombination in HAstV variation [25,27]. In agreement with most reports, we found no clear evidence of recombination between the 192-BJ07 strain and other

HAstV genotypes based on similarity plot analysis. Diversification of the HAstV amino sequences may be attributed to accumulated single nucleotide mutations. This mechanism is similar to the antigen drift in other viruses, such as in influenza viruses [28,29], which could lead to HAstVs escaping from existing host immunities and could result in the emergence of a new epidemic HAstV strain [30]. Additional studies, such as large scale whole genome sequencing, are needed to address the evolutionary patterns of HAstVs.

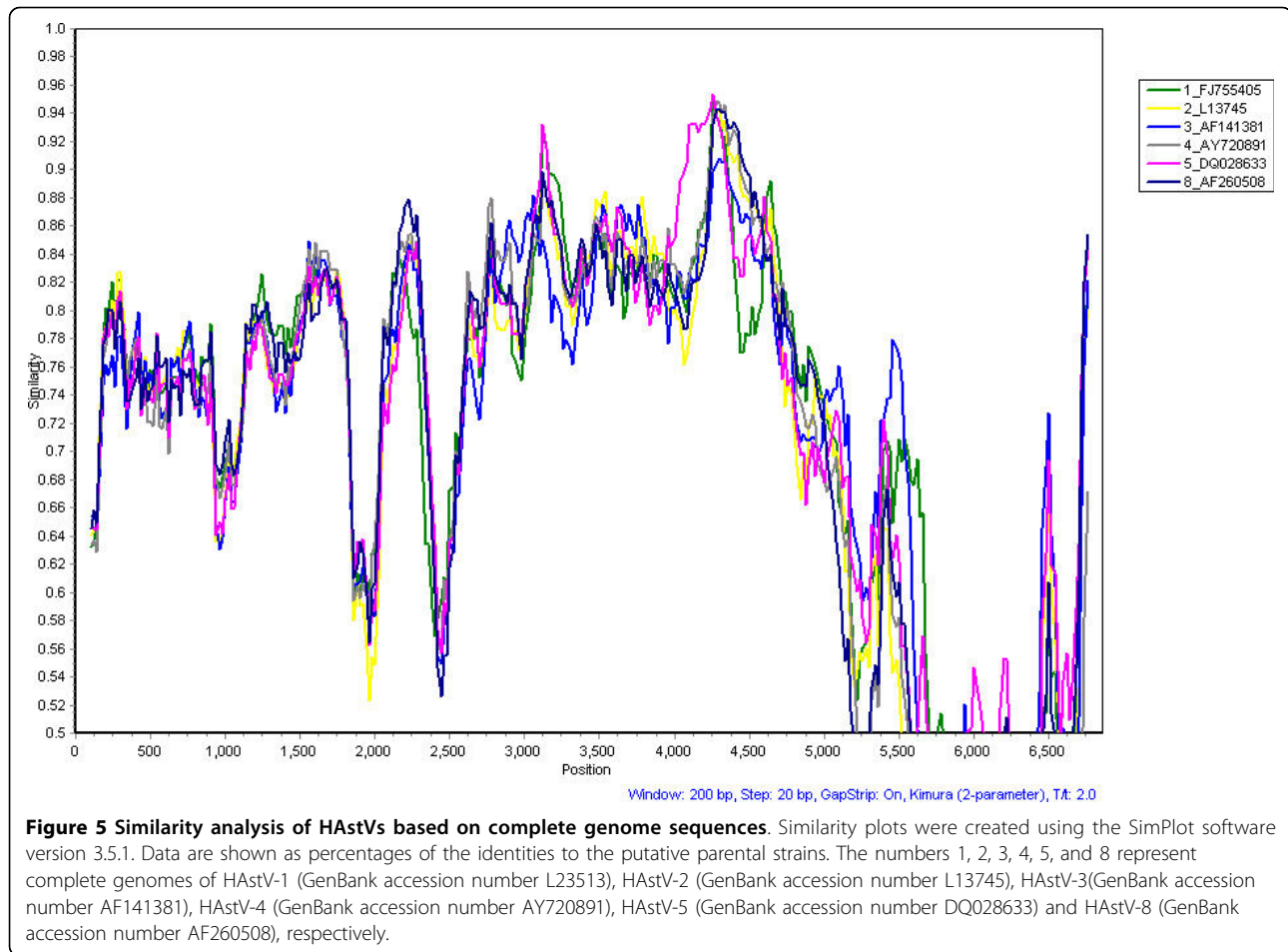
Conclusion

We have sequenced and characterized the complete genome of HAstV-6 (192-BJ07). This sequence will provide insight into the genetics of astroviruses, broaden our understanding of their properties, and inform surveillance and control of HAstV gastroenteritis around the world.

Methods

RNA extraction

A stool sample (termed 192-BJ07) that tested positive for HAstV-6 by RT-PCR was collected from a 2-year old boy who visited the Beijing Children's Hospital with acute diarrhea in 2007 [18]. Viral RNA was extracted



from the stool supernatant using Trizol reagent (Invitrogen, Carlsbad, CA) according to the manufacturer's instructions.

ORF2 amplification

The primers ORF2-F (5'-atggctagcaagctgacaagcagg-3') and ORF2-R (5'-gaagctgtaccctcgatcctactc-3') targeting ORF2 of 192-BJ07 were designed based on the only available HAstV-6 sequence in GenBank (GenBank accession number Z46658). For reverse transcription (RT) reactions, cDNA was generated with the SuperScript™ III RT kit (Invitrogen, Carlsbad, CA) using a random primer (Takara, Dalian, China) as described in the manufacturer's protocol. The PCR reaction was performed as follows: 94°C for 3 minutes, 35 cycles of amplification (94°C for 30 seconds; 50°C for 30 seconds; and 72°C for 3 minutes), and a final 10 minutes extension at 72°C. The PCR products were analyzed by 1.0% agarose gel electrophoresis and stained with ethidium bromide.

Genome amplification and sequencing

Rapid amplification of cDNA end (RACE) reactions were performed to obtain the entire sequence of the

viral genome by using the 5'- and 3'-RACE System for Rapid Amplification of cDNA Ends kit (Invitrogen, Carlsbad, CA) according to the manufacturer's protocol. The ORF2 sequence obtained above was used as the starting point for the amplification. PCR-amplified products were cloned into the pMD18-T vector (TaKaRa, Dalian, China) and were introduced into chemically competent *E. coli* DH5 α cells. The plasmid DNA was sequenced using an ABI3730 DNA Analyzer (Applied Biosystems). The complete genome sequence of HAstV-6 has been deposited in GenBank (GenBank Accession number GQ495608).

ORF prediction and RNA structure analysis

ORF1a and ORF2 were predicted for HAstV-6 192-BJ07 using the DNASTar ORF search program. ORF1b was predicted based on the "shifty" heptanucleotide (AAAAAAC) that occurs in other HAstVs [9]. RNA secondary structures were evaluated using RNAstructure 4.5 software.

Phylogenetic analysis

The MegAlign programs in the DNASTar software package were used to perform multiple sequence alignments.

HASV phylogenies with 1000 bootstrap replicates were created using the neighbor-joining method and the Kimura two-parameter model with the MEGA software version 4.0 [31].

Similarity analysis

SimPlot software version 3.5.1 [32] was used to analyze the relationships among the aligned HASV genome sequences. The complete genome sequences of 192-BJ07, HASV-1 (GenBank accession numbers L23513), HASV-2 (GenBank accession number L13745), HASV-3 (GenBank accession number AF141381), HASV-4 (GenBank accession numbers AY720891), HASV-5 (GenBank accession number DQ028633), and HASV-8 (GenBank accession number AF260508) were first aligned by using Clustal W of the MEGA 4 program, and then 192-BJ07 was chosen as the query sequence for the similarity analysis. Similarity was calculated in each window of 200 bp using the Kimura two-parameter method.

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Authors' contributions

JW, RG, GPB, and GV conceived the study. LG and JW designed the experiments. LG and WW carried out the experiments and analysis. YL participated in sequence analysis. LG, RG, and JW wrote the manuscript. All authors critically read and approved the final manuscript.

Competing interests

The authors declare that they have no competing interests.

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References

1. Chen SY, Chang YC, Lee YS, Chao HC, Tsao KC, Lin TY, Ko TY, Tsai CN, Chiu CH: **Molecular epidemiology and clinical manifestations of viral gastroenteritis in hospitalized pediatric patients in Northern Taiwan.** *J Clin Microbiol* 2007, **45**:2054-2057.
2. Guix S, Caballero S, Villena C, Bartolomé R, Latorre C, Rabella N, Simó M, Bosch A, Pintó RM: **Molecular epidemiology of astrovirus infection in Barcelona, Spain.** *J Clin Microbiol* 2002, **40**:133-139.
3. Liu MQ, Yang BF, Peng JS, Zhou DJ, Tang L, Wang B, Liu Y, Sun SH, Ho WZ: **Molecular epidemiology of astrovirus infection in infants in Wuhan, China.** *J Clin Microbiol* 2007, **45**:1308-1309.
4. Madeley CR, Cosgrove BP: **Letter: Viruses in infantile gastroenteritis.** *Lancet* 1975, **2**:124.
5. Mendez E, Arias CF: **Astroviruses.** *Fields Virology* Philadelphia: Lippincott Williams & Wilkins/Knipe DM, Howley PM, 5 2007, **1**:981-1000.
6. Kurtz JB, Lee TW: **Human astrovirus serotypes.** *Lancet* 1984, **2**:1405.
7. Koopmans MP, Bijen MH, Monroe SS, Vinjé J: **Age-stratified seroprevalence of neutralizing antibodies to astrovirus types 1 to 7 in humans in The Netherlands.** *Clin Diagn Lab Immunol* 1998, **5**:33-37.
8. Noel JS, Lee TW, Kurtz JB, Glass RI, Monroe SS: **Typing of human astroviruses from clinical isolates by enzyme immunoassay and nucleotide sequencing.** *J Clin Microbiol* 1995, **33**:797-801.
9. Jiang B, Monroe SS, Koonin EV, Stine SE, Glass RI: **RNA sequence of astrovirus: distinctive genomic organization and a putative retrovirus-like ribosomal frameshifting signal that directs the viral replicase synthesis.** *Proc Natl Acad Sci USA* 1993, **90**:10539-10543.
10. Lewis TL, Greenberg HB, Herrmann JE, Smith LS, Matsui SM: **Analysis of astrovirus serotype 1 RNA, identification of the viral RNA-dependent RNA polymerase motif, and expression of a viral structural protein.** *J Virol* 1994, **68**:77-83.
11. Oh D, Schreier E: **Molecular characterization of human astroviruses in Germany.** *Arch Virol* 2001, **146**:443-455.
12. Silva PA, Cardoso DD, Schreier E: **Molecular characterization of human astroviruses isolated in Brazil, including the complete sequences of astrovirus genotypes 4 and 5.** *Arch Virol* 2006, **151**:1405-1417.
13. Glass RI, Noel J, Mitchell D, Herrmann JE, Blacklow NR, Pickering LK, Dennehy P, Ruiz-Palacios G, de Guerrero ML, Monroe SS: **The changing epidemiology of astrovirus-associated gastroenteritis: a review.** *Arch Virol Suppl* 1996, **12**:287-300.
14. Lee TW, Kurtz JB: **Prevalence of human astrovirus serotypes in the Oxford region 1976-92, with evidence for two new serotypes.** *Epidemiol Infect* 1994, **112**:187-193.
15. Sakon N, Yamazaki K, Utagawa E, Okuno Y, Oishi I: **Genomic characterization of human astrovirus type 6 Katano virus and the establishment of a rapid and effective reverse transcription-polymerase chain reaction to detect all serotypes of human astrovirus.** *J Med Virol* 2000, **61**:125-131.
16. Gabbay YB, Linhares AC, Cavalcante-Pepino EL, Nakamura LS, Oliveira DS, da Silva LD, Mascarenhas JD, Oliveira CS, Monteiro TA, Leite JP: **Prevalence of human astrovirus genotypes associated with acute gastroenteritis among children in Belém, Brazil.** *J Med Virol* 2007, **79**:530-538.
17. Oishi I, Yamazaki K, Kimoto T, Minekawa Y, Utagawa E, Yamazaki S, Inouye S, Grohmann GS, Monroe SS, Stine SE: **A large outbreak of acute gastroenteritis associated with astrovirus among students and teachers in Osaka, Japan.** *J Infect Dis* 1994, **170**:439-443.
18. Guo L, Xu X, Song J, Wang W, Wang J, Hung T: **Molecular characterization of astrovirus infection in children with diarrhea in Beijing, 2005-2007.** *J Med Virol* 2010, **82**:415-423.
19. Bass DM, Qiu S: **Proteolytic processing of the astrovirus capsid.** *J Virol* 2000, **74**:1810-1814.
20. Méndez-Toss M, Romero-Guido P, Munguía ME, Méndez E, Arias CF: **Molecular analysis of a serotype 8 human astrovirus genome.** *J Gen Virol* 2000, **81**:2891-2897.
21. Willcocks MM, Kurtz JB, Lee TW, Carter MJ: **Prevalence of human astrovirus serotype 4: capsid protein sequence and comparison with other strains.** *Epidemiol Infect* 1995, **114**:385-391.
22. Lewis TL, Matsui SM: **Astrovirus ribosomal frameshifting in an infection-transfection transient expression system.** *J Virol* 1996, **70**:2869-2875.
23. Finkbeiner SR, Kirkwood CD, Wang D: **Complete genome sequence of a highly divergent astrovirus isolated from a child with acutediarrhea.** *Virology* 2008, **5**:117.
24. Walter JE, Briggs J, Guerrero ML, Matson DO, Pickering LK, Ruiz-Palacios G, Berke T, Mitchell DK: **Molecular characterization of a novel recombinant strain of human astrovirus associated with gastroenteritis in children.** *Arch Virol* 2001, **146**:2357-2367.
25. Lukashov VV, Goudsmit J: **Evolutionary relationships among Astroviridae.** *J Gen Virol* 2002, **83**:1397-1405.
26. Willcocks MM, Brown TD, Madeley CR, Carter MJ: **The complete sequence of a human astrovirus.** *J Gen Virol* 1994, **75**:1785-1788.
27. Ulloa JC, Matiz A, Lareo L, Gutierrez MF: **Molecular analysis of a 348 base-pair segment of open reading frame 2 of human astrovirus. A characterization of Colombian isolates.** *In Silico Biol* 2005, **5**:537-546.
28. Blackburne BP, Hay AJ, Goldstein RA: **Changing selective pressure during antigenic changes in human influenza H3.** *PLoS Pathog* 2008, **4**:e1000058.
29. Shen J, Ma J, Wang Q: **Evolutionary trends of A(H1N1) influenza virus hemagglutinin since 1918.** *PLoS One* 2009, **4**:e7789.
30. Tu ET, Bull RA, Greening GE, Hewitt J, Lyon MJ, Marshall JA, McIver CJ, Rawlinson WD, White PA: **Epidemics of gastroenteritis during 2006 were associated with the spread of norovirus GII.4 variants 2006a and 2006b.** *Clin Infect Dis* 2008, **46**:413-420.

31. Tamura K, Dudley J, Nei M, Kumar S: **MEGA4: Molecular Evolutionary Genetics Analysis (MEGA) software version 4.0.** *Mol Biol Evol* 2007, **24**:1596-1599.
32. Lole KS, Bollinger RC, Paranjape RS, Gadkari D, Kulkarni SS, Novak NG, Ingersoll R, Sheppard HW, Ray SC: **Full-length human immunodeficiency virus type 1 genomes from subtype C-infected seroconverters in India, with evidence of intersubtype recombination.** *J Virol* 1999, **73**:152-160.

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