**Antigenic Diversity of Human Sapoviruses**

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Sapovirus (SaV) is a causative agent of gastroenteritis. On the basis of capsid protein (VP1) nucleotide sequences, SaV can be divided into 5 genogroups (GI–GV), of which the GI, GII, GIV, and GV strains infect humans. SaV is unculivable, but expression of recombinant VP1 in insect cells results in formation of viruslike particles (VLPs) that are antigenically similar to native SaV. In this study, we newly expressed SaV GII and GIV VLPs to compare genetic and antigenic relationships among all human SaV genogroups. Hyperimmune antiserum samples against VLPs reacted strongly with homologous VLPs. However, several antiserum samples weakly cross-reacted against heterologous VLPs in an antibody ELISA. Conversely, an antigen ELISA showed that VLPs of SaV in all human genogroups were antigenically distinct. These findings indicate a likely correspondence between SaV antigenicity and VP1 genogrouping and genotyping.

The family Caliciviridae contains 4 genera (Sapovirus, Norovirus, Lagovirus, and Vesivirus), which include Sapporo virus, Norwalk virus, Rabbit hemorrhagic disease virus, and Feline calicivirus, respectively. Sapoviruses (SaVs) and noroviruses (NoVs) are etiologic agents of human gastroenteritis. The prototype strain of human SaV, Sapporo virus, was originally discovered in an outbreak in an orphanage in Sapporo, Japan, in 1977 (1). SaV infects children and adults and has been found to cause outbreaks of gastroenteritis in daycare centers, healthcare facilities, and elementary schools. Detection methods include reverse transcription–PCR (RT-PCR), real-time RT-PCR, enzyme immunoassays, and ELISAs (2–6). Recently, we detected SaV in untreated wastewater samples, treated wastewater samples, and river samples (7).

The SaV genomes are predicted to contain either 2 or 3 main open reading frames (ORF1–3). SaV ORF1 encodes for nonstructural proteins and the major capsid protein, and ORF2 (VP2) and ORF3 encode proteins of yet unknown functions. On the basis of VP1 nucleotide sequences, SaVs have been divided into 5 genogroups (GI–GV), of which GI, GII, GIV, and GV strains infect humans and GIII strains infect porcine species (8). SaV genogroups can be further subdivided into genotypes. Recently, we identified several recombinant SaV strains (8,9).

Human SaV and NoV strains are uncultivable, but expression of a recombinant subgenomic-like construct (i.e., VP1 to the end of the genome) or VP1 alone in insect or mammalian cells results in the formation of viruslike particles (VLPs) that are morphologically similar to native SaV (10–16). However, production of VLPs of SaV remains difficult, usually only resulting in low yields of VLPs compared with norovirus (10,12,16,17). Cryoelectron microscopy and x-ray crystallography analyses of NoV VLPs identified the shell (S) and protruding domains (subdomains P1–1, P1–2, and P2) (18). Also, Chen et al. described strictly and moderately conserved amino acid residues in the capsid protein among the 4 genera in the family Caliciviridae (13).

Previously, we reported that SaV GI/1 (strain Mc114) and GV/1 (strain NK24) were antigenically distinct (5,10). More recently, we discovered that SaV GI/5 (strain Yokote1) VLPs were antigenically distinct from SaV GI/1 Mc114 and GV/1 NK24 VLPs (19). Other than these few studies, little is known about the genetic and antigenic relationships among the 4 human SaV genogroups. For classification of NoV, distinct genotypes have been defined as having bootstrap values >950 (VP1 sequences); at least 14 GI and 17 GII genotypes have been identified (20). For SaV, genogroups have only been vaguely defined, mostly

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because 2 of them (GIV/1 and GV/1) were only recently identified, few sequences exist in the database, and antigenic relationships among all genogroups are unknown. In addition, genetic recombination was only recently discovered and appears to be common within the genus Sapovirus.

The purpose of this study was to examine cross-reactivities among the 4 human SaV genogroups and compare results with those of genetic analysis. For this purpose, 2 other SaV strains, GI/3 Syd53 and GIV/1 Syd3, were expressed and antisera were produced against their purified VLPs. A total of 5 SaV strains (GI/1 Mc114, GI/5 Yokote1, GI/3 Syd53, GIV/1 Syd3, and GV/1 NK24) that include all 4 human genogroups and 2 GI genotypes were compared. Our results show that SaV genogroups were antigenically distinct and corresponded with results of genetic classification on the basis of full-length VP1 nucleotide sequences. Proper genetic classification of SaV strains is required, and a consensus of genogroups and genotypes that represent genetically and antigenically diverse strains, which include recombinant SaV strains, should be established to avoid conflicting grouping.

Materials and Methods

Specimens

Virus-positive stool specimens were collected from several sources. SaV strain Mc114 (GenBank accession no. AY237422) was isolated from an infant hospitalized with acute gastroenteritis in Chiang Mai, Thailand, in 2001 (21). SaV strain NK24 (AY646856) was isolated from an infant with gastroenteritis in Nong Khai, Thailand, in 2003 (22). SaV strain Yokote1 was isolated from an outbreak of gastroenteritis at a kindergarten in Yokote City, Japan, in 2006 (19). SaV strains Syd53 and Syd3 were isolated from infants hospitalized with acute gastroenteritis in Sydney, New South Wales, Australia, in 2001 (23). NoV strain Osaka659 was isolated from an outbreak of gastroenteritis in Japan, in 2006 (unpub. data). RNA extraction and RT-PCR were performed as described (24).

Sequence Analysis

Nucleotide sequences were determined by using the Terminator Cycle Sequence Kit version 3.1 and an ABI 3130 sequencer (both from Applied Biosystems, Boston, MA, USA). Nucleotide sequences were aligned with ClustalX (www.ebi.ac.uk/clustal/darwin) and the distances were calculated by the Kimura 2-parameter method (24). Phylogenetic trees with bootstrap analysis from 1,000 replicates were generated by the neighbor-joining method as described (20). Amino acid VP1 secondary structure predictions were made by using PSIPRED secondary structural prediction software (25).

Expression of Viralike Particles

For the expression of VP1 in insect cells, all SaV constructs were designed to begin from the predicted VP1 start AUG codon and included the ORF2 and poly(A) sequences. SaV strains Syd53 and Syd3 were cloned as described (10) for strains SaV Mc114, NK24, and Yokote1 according to the protocol of the Baculovirus Expression System using Gateway Technology (Invitrogen, Carlsbad, CA, USA). Briefly, strains Syd53 and Syd3 were amplified with specific sense primers Syd53attb1 (5'-GGGGACAAAGTTGGTCAAAAAACAGAGGCTTCAAGAGATAGAACCATGGAGGGTGTTGTCCCACCAGA-3') and Syd3attb1 (5'-GAGGAGGGTGTTGCTCCACCAGA-3') and further amplified with specific antisense primer Syd53attb2 (5'-GAGGAGGGTGTTGCTCCACCAGA-3') and Syd3attb2 (5'-GAGGAGGGTGTTGCTCCACCAGA-3'). PCR fragments were purified after electrophoresis on a 1.0% agarose gel. Fragments were cloned into donor vector pDONR201 (Invitrogen) and transferred into a baculovirus transfer vector pDEST8 (Invitrogen).

The recombinant pDEST8 was purified and used to transform Escherichia coli DH10Bac-competent cells (Invitrogen), which produced recombinant bacmids (baculovirus shuttle vectors) containing the VP1 gene. Recombinant bacmids were then transfected into Sf9 cells (Riken Cell Bank, Ibaraki, Japan), and recombinant baculoviruses were isolated. Recombinant baculoviruses were used to infect ≈3 × 10⁶ confluent Tn5 cells (Invitrogen) at a multiplicity of infection of 5–10 in 1.5 mL of Ex-Cell 405 medium (JRH Biosciences, Lenexa, KS, USA), and the infected cells were incubated at 26°C. The culture medium was harvested 5–6 d postinfection, centrifuged for 10 min at 3,000× g, and further centrifuged for 30 min at 10,000× g. VLPs were concentrated by ultracentrifugation for 2 h at 45,000 rpm at 4°C (Beckman TLA-55 rotor; Beckman Coulter, Fullerton, CA, USA), and resuspended in 30 μL of Grace’s medium. Samples were examined for VLP formation by electron microscopy as described (10), and large-scale production of VLPs was performed as described (24).

Antibody Production

Hyperimmune sera to newly expressed VLPs of SaV (Syd53 and Syd3) were prepared in rabbits and guinea pigs. The first subcutaneous injection was performed with purified VLPs (≈10 μg) in Freund complete adjuvant. After 3 weeks, the animals received 1 booster injection (intravenously in rabbits and subcutaneously in guinea pigs) of 10 μg of VLPs without adjuvant. Blood was collected from the animals 1 week after their last booster injection.
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**Antibody ELISA**

Cross-reactivities among antiserum samples against SaV were examined by using an antibody ELISA with hyperimmune rabbit antibodies against VLPs. Briefly, wells of 96-well microtiter plates (Maxisorp; Nunc, Roskilde, Denmark) were each coated with 100 μL of purified VLPs (≈1.0 μg/mL in carbonate-bicarbonate buffer, pH 9.6) (Sigma, St. Louis, MO, USA) and incubated overnight at 4°C. Wells were washed twice with phosphate-buffered saline (PBS) containing 0.1% Tween 20 (PBS-T) and blocked with PBS containing 5% skim milk (PBS-SM) for 1 h at room temperature. Wells were then washed 4 times with PBS-T, 100 μL of 2-fold–diluted hyperimmune rabbit antibodies from an initial concentration of 1:500 in PBS-T-SM was added to each well, and the plates were incubated for 1 h at 37°C. Wells were then washed 4 times with PBS-T, and 100 μL of a 1:1,000 dilution of horseradish peroxidase–conjugated goat antirabbit immunoglobulin G diluted in PBS-T-SM was added to each well. Plates were incubated for 1 h at 37°C. Wells were then washed 4 times with PBS-T, and 100 μL of substrate (o-phenylenediamine) and H2O2 were added to each well, and the plates were left in the dark for 30 min at room temperature. The reaction was stopped by the addition of 50 μL of 2N H2SO4 to each well, and the absorbance was measured at 492 nm (A492). The optical density (OD) cutoff point was determined to be 0.15, which was equal to 3 times the mean OD of preimmune serum (5).

**Antigen ELISA**

Cross-reactivities among VLPs were also examined by using an antigen ELISA. Briefly, wells were coated with 100 μL of a 1:8,000 dilution of hyperimmune rabbit antiserum diluted in PBS (except for Syd3, for which a 1:3,000 dilution was used), and the plates were incubated overnight at 4°C. Wells were washed 4 times with PBS-T and blocked with PBS-SM for 1 h at room temperature. Wells were then washed 4 times with PBS-T, 100 μL of VLPs (≈1.0 μg/mL in carbonate-bicarbonate buffer, pH 9.6) (Sigma) was added to duplicate hyperimmune rabbit wells, and the plates were incubated for 1 h at 37°C. Wells were then washed 4 times with PBS-T, 100 μL of a 1:1,000 dilution of hyperimmune pig antibody diluted in PBS-T-SM was added to each well (except for Syd3, which used a 1:3,000 dilution), and the plates were incubated for 1 h at 37°C. Wells were washed 4 times with PBS-T, and 100 μL of a 1:1,000 dilution of horseradish peroxidase–conjugated rabbit antiguinea pig immunoglobulin G diluted in PBS-T-SM was added to each well. Plates were then processed as described above. On the basis of our previous study, a specimen with an A492 (P – N) >0.1 and a P/N ratio >1.34 (where P is hyperimmune antiserum and N is preimmune antiserum) was considered significantly positive (4).

**Results**

**Sequence Analysis**

The sequence of the 3’ end of the genome (≈2,600 nt), i.e., VP1 to poly(A), for the newly expressed SaV strains (Syd53 and Syd3) was determined. Genetic analysis was performed with only complete VP1 sequences, which included sequences from our epidemiologic studies and other sequences available on the database (Figure 1). Five SaV GI and 6 GII genotypes were observed, but only 1 genotype for SaV GIV and 1 for GV was found. This result suggests that SaV GI and GII strains were more genetically diverse, prevalent, or more virulent than SaV GIV and GV strains. However, because the SaV GIV and GV strains were only recently detected (26,27), this result may reflect only the specificity and sensitivity of the detection methods used. On the basis of our previous classifications, SaV Mc114 and Yokote1 sequences both belonged to GI, but to different genotypes, GI/1 and GI/5, respectively; Syd53 be-

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**Figure 1. Phylogenetic tree of sapoviruses on the basis of entire nucleotide sequences of the capsid protein.** Different genogroups and genotypes are indicated. The numbers on each branch indicate the bootstrap values for the genotype. The scale bar at the lower left represents nucleotide substitutions per site. GenBank accession nos. for reference strains: Arg39, AY289803; Bristol, HCA249939; C12, AY603425; Chiba000496, AJ412800; Chiba10658, AY606696; Cruise ship, AY289804; Dresden, AY694184; Ehime643, DQ366345; Ehime1107, DQ058829; Houston27, U95644; Manchester, X86560; Mc2, AY237419; Mc10, AY237420; Mc114, AY237422; Mex340, AF435812; NK24, AY646856; Parkville, U73124; PEC, AF182760; Potsdam, AF294739; Sapporo, U65427; SK15, AY648855; Stockholm, AF194182; SW278, DQ125333; Syd3, DQ104357; Syd53, DQ104360; and Yokote1, AB253740.

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longed to GII/3; Syd3 belonged to GIV/1; and NK24 belonged to GV/1. SaV GI/1 Mc114 and GI/5 Yokote1 VP1 sequences shared 76.5% and 79% nucleotide similarity and amino acid identity, respectively (Table 1). The nucleotide similarity and amino acid identity among the genogroups was low, i.e., <60% (Table 1).

Expression of VP1

We previously expressed SaV GI/1 Mc114, GI/5 Yokote1, and GV/1 NK24 in insect cells, which resulted in the formation of VLPs morphologically similar to native SaV (5,10). In this study, we newly expressed SaV GII/3 Syd53 and GIV/1 Syd3 in insect cells to analyze the cross-reactivity among all human SaV genogroups. SaV GII/3 Syd53 and GIV/1 Syd3 successfully formed VLPs with a diameter of 41 to 46 nm and were morphologically similar to native SaV (Figure 2). Hyperimmune sera against these purified VLPs were prepared in rabbits and guinea pigs.

Antibody ELISA Analysis

Our previous antibody ELISA result showed that SaV GI/1 Mc114 and GV/1 NK24 antiserum had no cross-reactivities against heterologous GV/1 NK24 and GI/1 Mc114 VLPs, respectively (5). In the current study, cross-reactivities among 5 VLPs of SaV (GI/1 Mc114, GI/5 Yokote1, GII/3 Syd53, GIV/1 Syd3, and GV/1 NK24) were analyzed by using the antibody ELISA with 2-fold serial dilutions (1:500–1:1,024,000) of hyperimmune antiserum (Figure 3). The OD cutoff point was 0.15, which was equal to 3 times the mean OD of preimmune serum (5). Hyperimmune rabbit antiserum reacted strongly against the homologous VLPs (Table 2, Figure 3). SaV GII/3 Syd53 and GIV/1 Syd3 antiserum titers were 512,000 and 2,056,000, respectively (Table 2). Several antiserum weakly cross-reacted with heterologous VLPs. SaV GI/1 Mc114 antiserum cross-reacted weakly with GI/5 Yokote1 and GII/3 Syd53 VLPs, i.e., their cross-reactivities were 8- and 16-fold lower than that of the homologous VLP titer, respectively. SaV GI/5 Yokote1 antiserum cross-reacted weakly with GI/1 Mc114 and GII/3 Syd53 VLPs, i.e., their cross-reactivity was 16-fold lower than that of the homologous VLP titer. SaV GII/3 Syd53, GIV/1 Syd3, and GV/1 NK24 antiserum appeared to have no cross-reactivities against any of the heterologous VLPs, i.e., their cross-reactivities were >32-fold lower than those of the homologous VLP titer. These results suggested that SaV GI/1 Mc114 and GI/5 Yokote1 antiserum had weak 2-way cross-reactivities against GII/3 Syd53, GIV/1 Syd3, and GV/1 NK24 VLPs, respectively. The negative control NoV Osaka659 antiserum showed no cross-reactivities against VLPs of SaV at any dilution of antiserum, which indicates that the antiserum was specific for the VLPs and not the insect cell proteins.

Antigen ELISA Analysis

On the basis of a previous study, a specimen with an $A_{492} (P – N) >0.10$ and a P/N ratio >1.34 was considered significantly positive (4). Our recent antigen ELISA results showed that SaV GI Mc114, GI/5 Yokote1, and GV/1 NK24 VLPs were antigenically distinct (19), i.e., GI/1 Mc114 $P–N 0.41$, P/N 9.19; GI/5 Yokote1 $P – N 0.93$.
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P/N 19.59; and GV/1 NK24 P – N 1.03, P/N 21.58. In the current study, we examined the cross-reactivity among the newly expressed VLPs and those previously prepared by using the antigen ELISA. The antiserum samples reacted only with homologous VLPs, i.e., SaV GII/3 Syd53 P – N 1.02, P/N 21.33; and GIV/1 Syd3 P – N 1.44, P/N 29.74 (Table 3). Several antiseras appeared to cross-react with heterologous VLPs, i.e., where the P – N was <0.10, but the P/N ratio was >1.34. SaV GI/5 Yokote1 antiserum appeared to cross-react with GII/3 Syd53 VLPs (P/N 1.97); GIV/1 Syd3 antiserum appeared to cross-react with GI/5 Yokote1 VLPs (P/N 1.42); and GV/1 NK24 antiserum appeared to cross-react with GII/3 Syd53 VLPs (P/N 1.50). However, all of these cross-reactivity results were considered negative because P – N was <0.10.

Amino Acid Alignment and Secondary Structure Prediction

An amino acid alignment of the 5 SaV VP1 sequences (GI/1 Mc114, GI/5 Yokote1, GII/3 Syd53, GIV Syd3, and GV NK24) showed that the shell domain contained more conserved residues than the predicted P domains (Figure 4). However, SaV GI/1 Mc114 and GI/5 Yokote1 shared more conserved continuous residues in the predicted P2 subdomain than other genogroups. The NoV P2 subdomain is thought to contain the determinants of strain specificity, cell binding, and antigenicity. For example, monoclonal antibodies that recognize regions in the P2 subdomain inhibit binding of NoV VLPs to cells (28,29). In a recent study, we analyzed cross-reactivities among 26 different NoV VLPs (6 GI and 12 GII genotypes) (30) and found

| Antiserum | Mc114 (GI/1) | Yokote1 (GI/5) | Syd53 (GII/3) | Syd3 (GIV/1) | NK24 (GV/1) |
|-----------|--------------|----------------|--------------|-------------|-------------|
| Mc114 (GI/1) | 512,000      | 64,000         | 32,000       | 1,000       | <1,000      |
| Yokote1 (GI/5) | 32,000      | 512,000        | 32,000       | 2,000       | 4,000       |
| Syd53 (GII/3) | 4,000        | 8,000          | 512,000      | 4,000       | <1,000      |
| Syd3 (GIV/1) | 2,000        | 16,000         | 8,000        | 2,056,000   | 1,000       |
| NK24 (GV/1) | 2,000        | 2,000          | 4,000        | 1,000       | 2,056,000   |

Figure 3. Antibody ELISA absorbances at 492 nm of 6 viruslike particles (VLPs) of sapovirus. A) Mc114 GI/1; B) Yokote1 GI/5; C) Syd53 GII/3; D) Syd3 GIV/1; E) NK24 GV/1; F) NoV Osaka659. Wells were coated with =100 ng of purified VLPs. Antiserum was used in 2-fold dilutions from 1:500 to 1,024,000 in phosphate-buffered saline, 0.1% Tween 20, 5% skim milk shown as 1–12 along the x-axis. OD, optical density; NoV, Norwalk virus.

Table 2. Titers of antiseras to viruslike particles of sapovirus strains in an antibody ELISA

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broad-range cross-reactivities for several NoV antisera. Our results suggested that these cross-reactivities were due to conserved amino acid residues located outside the P2 domain. Conversely, secondary structure predictions made by using PSIPRED secondary structural prediction software showed that helix structures could also influence the cross-reactivity among the NoV VLPs. In the current study, we determined the secondary structure of the 5 SaV VP1 amino acid sequences. Overall, SaV VP1 structures appear to be similar (online Appendix Figure, available from www.cdc.gov/EID/content/13/10/1519-appG.htm). The location of 3 helix structures in the shell domain and 1 helix structure in the C-terminal region were nearly identical for the 5 SaV VP1 sequences. Only SaV GV/1 NK24 was predicted to have a single helix structure in the P2 subdomain. These results suggested that the amino acid sequence, particularly the P2 subdomain, plays a major role in determining cross-reactivity among SaV strains. However, additional studies, including high-resolution VLP structural analysis, are needed.

Discussion

In this study, we analyzed genetic and antigenic relationships for 4 human SaV genogroups (GI, GII, GIV, and GV). We observed weak 2-way cross-reactivity with SaV GI/1 Mc114 and GI/5 Yokote1 antisera against the heterologous GI/5 Yokote1 and GI/1 Mc114 VLPs, respectively, by using an antibody ELISA. However, when we used an antigen ELISA, we found that GI/1 Mc114 and GI/5 Yokote1 VLPs were antigenically distinct. These weak cross-reactivities identified by using the antibody ELISA may have been influenced by several factors, including unfolded VLPs on the microtiter plates at the high pH (carbonate-bicarbonate buffer, pH 9.6) (31) or conserved continuous residues outside the predicted P2 domain. Therefore, these 2 SaV genotypes (GI/1 and GI/5) are for the most part antigenically distinct. Likewise, we found that the 4 human SaV genogroups were antigenically distinct in the antigen ELISA. To our knowledge, these new findings provide the first evidence that SaV antigenicity corresponded well with VP1 genotyping and genotyping.
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Dr Hansman is a scientist at the National Institute of Infectious Diseases, Tokyo, Japan. His research interests include the epidemiology, expression, and cross-reactivity of viruses that cause gastroenteritis in humans, namely, sapovirus and norovirus.

References

1. Chiba S, Sakuma Y, Kogasaka R, Akihara M, Horino K, Nakao T, et al. An outbreak of gastroenteritis associated with calcivirus in an infant home. J Med Virol. 1979;4:249–54.

2. Oka T, Katayama K, Hansman GS, Kageyama T, Ogawa S, Wu FT, et al. Detection of human sapovirus by real-time reverse transcription-polymerase chain reaction. J Med Virol. 2006;78:1347–53.

3. Nakata S, Chiba S, Terashima H, Sakuma Y, Kogasaka R, Nakao T. Microtiter solid-phase radioimmunoasay for detection of human calicivirus in stools. J Clin Microbiol. 1983;17:198–201.

4. Hansman GS, Guntapon R, Pongsuwanma Y, Natori K, Katayama K, Takeda N. Development of an antigen ELISA to detect sapovirus in clinical stool specimens. Arch Virol. 2006;151:551–61.

5. Hansman GS, Natori K, Ushijima H, Katayama K, Takeda N. Characterization of polyclonal antibodies raised against sapovirus genogroup five virus-like particles. Arch Virol. 2005;150:1433–7.

6. Okada M, Yamashita Y, Osato M, Shinozaki K. The detection of human sapoviruses with universal and genogroup-specific primers. Arch Virol. 2006;151:2503–9.

7. Hansman GS, Sano D, Ueki Y, Imai T, Oka T, Katayama K, et al. Sapovirus in water. Japan. Emerg Infect Dis. 2007;13:133–5.

8. Hansman GS, Takeda N, Oka T, Osato M, Hedlund KO, Katayama K. Intergenogroup recombination in sapoviruses. Emerg Infect Dis. 2005;11:1916–20.

9. Katayama K, Miyoshi T, Uchino K, Oka T, Tanaka T, Takeda N, et al. Novel recombinant sapovirus. Emerg Infect Dis. 2004;10:1874–6.

10. Hansman GS, Natori K, Oka T, Ogawa S, Tanaka K, Nagata N, et al. Cross-reactivity among sapovirus recombinant capsid proteins. Arch Virol. 2005;150:21–36.

11. Guo M, Qian Y, Chang KO, Saif LJ. Expression and self-assembly in baculovirus of porcine enteric calicivirus capsids into virus-like particles and their use in an enzyme-linked immunosorbent assay for antibody detection in swine. J Clin Microbiol. 2001;39:1487–93.

12. Hansman GS, Katayama K, Oka T, Natori K, Takeda N. Mutational study of sapovirus expression in insect cells. Virol J. 2005;2:13.

13. Chen R, Neill JD, Noel JS, Hutson AM, Glass RI, Estes MK, et al. Inter- and intragenus structural variations in caliciviruses and their functional implications. J Virol. 2004;78:6469–79.

14. Jiang X, Zhong W, Kaplan M, Pickering LK, Matson DO. Expression and characterization of Sapporo-like human calicivirus capsid proteins in baculovirus. J Virol Methods. 1999;78:81–91.

15. Numata K, Hardy ME, Nakata S, Chiba S, Estes MK. Molecular characterization of morphologically typical human calicivirus Sapporo. Arch Virol. 1997;142:1537–52.

16. Oka T, Hansman GS, Katayama K, Ogawa S, Nagata N, Miyamura T, et al. Expression of sapovirus virus-like particles in mammalian cells. Arch Virol. 2006;151:399–404.

17. Farkas T, Deng X, Ruiz-Palacios G, Morrow A, Jiang X. Development of an enzyme immunoassay for detection of sapovirus-specific antibodies and its application in a study of seroprevalence in children. J Clin Microbiol. 2006;44:3674–9.

18. Prasad BV, Hardy ME, Dokland T, Bella J, Rossmann MG, Estes MK. X-ray crystallographic structure of the Norwalk virus capsid. Science. 1999;286:287–90.

19. Hansman GS, Saito H, Shibata C, Ishizuka S, Osato M, Oka T, et al. Outbreak of gastroenteritis due to sapovirus. J Clin Microbiol. 2007;45:1347–9.

20. Kageyama T, Shinohara M, Uchida K, Fukushima S, Hoshino FB, Kojima S, et al. Coexistence of multiple genotypes, including newly identified genotypes, in outbreaks of gastroenteritis due to norovirus in Japan. J Clin Microbiol. 2004;42:2988–95.

21. Hansman GS, Katayama K, Maneekarn N, Peerakome S, Khamrin T, Tonusin S, et al. Genetic diversity of norovirus and sapovirus in hospitalized infants with sporadic cases of acute gastroenteritis in Chiang Mai, Thailand. J Clin Microbiol. 2004;42:1305–7.

22. Guntapon R, Hansman GS, Oka T, Ogawa S, Kageyama T, Pongsuwanma Y, et al. Norovirus and sapovirus infections in Thailand. Jpn J Infect Dis. 2004;57:276–8.

23. Hansman GS, Takeda N, Katayama K, Tu ET, McIver CJ, Rawlinson WD, et al. Genetic diversity of sapovirus in children, Australia. Emerg Infect Dis. 2006;12:141–3.

24. Katayama K, Shirato-Horikoshi H, Kojima S, Kageyama T, Oka T, Hoshino F, et al. Phylogenetic analysis of the complete genome of 18 Norwalk-like viruses. Virology. 2002;299:225–39.

25. McGuffin LJ, Bryson K, Jones DT. The PSIPRED protein structure prediction server. Bioinformatics. 2000;16:404–5.

26. Farkas T, Zhong WM, Jing Y, Huang PW, Espinosa SM, Martinez N, et al. Genetic diversity among sapoviruses. Arch Virol. 2004;149:1309–23.

27. Okada M, Shinozaki K, Ogawa T, Kaino I. Molecular epidemiology and phylogenetic analysis of Sapporo-like viruses. Arch Virol. 2002;147:1445–51.

28. Lochridge VP, Jutila KL, Graff JW, Hardy ME. Epitopes in the P2 domain of norovirus VP1 recognized by monoclonal antibodies that block cell interactions. J Gen Virol. 2005;86:2799–806.

29. White LJ, Ball JM, Hardy ME, Tanaka TN, Kitamoto N, Estes MK. Attachment and entry of recombinant Norwalk virus capsids to cultured human and animal cell lines. J Virol. 1996;70:6589–97.

30. Hansman GS, Natori K, Shirato-Horikoshi H, Kojima S, Kageyama T, Oka T, Hoshino F, et al. Phylogenetic analysis of the complete genome of 18 Norwalk-like viruses. Virology. 2002;299:225–39.

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