

Cross-reactivity among sapovirus recombinant capsid proteins

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Received May 14, 2004; accepted August 2, 2004

Published online September 24, 2004 © Springer-Verlag 2004

Summary. Sapovirus (SaV), a member of the genus *Sapovirus* in the family *Caliciviridae*, is an agent of human and porcine gastroenteritis. SaV strains are divided into five genogroups (GI–GV) based on their capsid (VP1) sequences. Human SaV strains are noncultivable, but expression of the recombinant capsid protein (rVP1) in a baculovirus expression system results in the self-assembly of virus-like particles (VLPs) that are morphologically similar to native SaV. In this study, rVP1 constructs of SaV GI, GII, and GV strains were expressed in a baculovirus expression system. The structures of the GI, GII, and GV VLPs, with diameters of 41–48 nm, were morphologically similar to those of native SaV. However a fraction of GV VLPs were smaller, with diameters of 26–31 nm and spikes on the outline. This is the first report of GII and GV VLP formation and the first identification of small VLPs. To examine the cross-reactivities among GI, GII, and GV rVP1, hyperimmune rabbit antisera were raised against *Escherichia coli*-expressed GI, GII, and GV N- and C-terminal VP1. Western blotting showed the GI antisera cross-reacted with GV rVP1 but not GII rVP1; GII antisera cross-reacted weakly with GI rVP1 but did not cross-react with GV rVP1; and GV antisera reacted only with GV rVP1. Also, hyperimmune rabbit and guinea pig antisera raised against purified GI VLPs were used to examine the cross-reactivities among GI, GII, and GV VLPs by an antigen enzyme-linked immunosorbent assay (ELISA). The ELISA showed that the GI VLPs were antigenically distinct from GII and GV VLPs.

Introduction

Human sapovirus (SaV) is a member of the genus *Sapovirus* in the family *Caliciviridae*. The prototype strain of human SaV, the Sapporo virus, was originally discovered from an outbreak in an orphanage in Sapporo, Japan, in October 1977 [5]. Chiba et al. identified viruses with the typical animal calicivirus morphology, called the “Star of David” structure, by electron microscopy (EM). In addition, SaV strains typically feature a diameter of 41–48 nm, cup-shaped depressions, and ten spikes on the outline.

Human SaV strains infect all age groups and can cause outbreaks of gastroenteritis and sporadic infections requiring hospitalization [7, 13, 18, 19, 21, 27]. Several groups have purified native SaV particles from stool specimens and produced antisera against them for use in immunoassays, including immune-EM and enzyme-linked immunosorbent assays (ELISAs) [14, 18, 20, 25]. However, the most widely used method to detect SaV is reverse transcription-polymerase chain reaction (RT-PCR), which has a high sensitivity [23]. SaV strains were recently divided into five genogroups (GI–GV), of which GI, GII, GIV, and GV strains infect humans, while GIII strains infect porcine species [6]. The SaV GI, GIV, and GV genomes are predicted to each contain three main open reading frames (ORFs), whereas SaV GII and GIII genomes each have only two main ORFs [6, 8, 17, 24]. The SaV ORF1 encodes the nonstructural proteins and the major capsid protein (VP1), while ORF2 and ORF3 encode proteins of yet-unknown functions. Therefore, the VP1 may be produced by either of two pathways: (i) translated as part of ORF1 and then cleaved, or (ii) translated from subgenomic RNA, although the subgenomic RNA of human SaV has not yet been identified.

Human SaV strains are noncultivable, but expression of the SaV recombinant VP1 (rVP1) in a baculovirus expression system results in the self-assembly of virus-like particles (VLPs) that are morphologically similar to native SaV [22]. There are four reports to date of the successful assembly of SaV VLPs [4, 9, 14, 22]. In three of these reports, however, the formation of VLPs was observed in rVP1 constructs that included short sequences upstream from the predicted rVP1 start AUG codon. Jiang et al. found that an upstream sequence of 73 nucleotides from the predicted VP1 start AUG codon was crucial for VLP formation [14], whereas the authors in two of the other three reports included 9 and 39 nucleotides upstream, respectively [9, 22]. The predicted human SaV VP1 start contains an amino acid motif, MEG, which is conserved in all human SaV strains and as such is considered the putative VP1 start. Recently, structural analysis of SaV VLPs predicted the shell (S) and protruding domains (subdomains P1 and P2) [4].

In the current study, we report the self-assembly of SaV GI, GII, and GV VLPs using constructs that began exactly from the predicted VP1 start AUG codon in a baculovirus expression system. More importantly, we describe for the first time the cross-reactivities among SaV GI, GII, and GV rVP1 by Western blotting and the cross-reactivities among SaV GI, GII, and GV VLPs by an antigen ELISA.

Materials and methods

Viruses

SaV Mc114 strain (GenBank accession number AY237422) was isolated from an infant hospitalized with acute gastroenteritis in Chiang Mai, Thailand, in 2001 [11]. SaV C12 strain (AY603425) was isolated from an infant with gastroenteritis in Sakai, Japan, in 2001 (in press). SaV NK24 strain (AY646856) was isolated from an infant with gastroenteritis in Nong Khai, Thailand, in 2003 (manuscript in press).

RT-PCR, sequencing, phylogenetic analysis

RNA extraction, RT-PCR, sequencing, and phylogenetic analysis were performed as previously described [15]. SaV sequences were phylogenetically classified based on the scheme of Farkas et al. [6].

Cloning of viral cDNA to produce recombinant bacmids

For the expression of rVP1 in insect cells, SaV constructs were designed to begin from the predicted VP1 start AUG codon, and included the VP2 and poly(A) sequences. PCR-amplified fragments were cloned according to the protocol of the Baculovirus Expression system using Gateway Technology (Invitrogen, USA). For the Mc114 strain, primers p+1Mc114 and attB2TX30SXN were used. For the C12 strain, primers p+1C12 and attB2TX30SXN were used. For the NK24 strain, primers p+1NK24 and attB2TX30SXN were used. PCR fragments were cut and purified from a 0.8% agarose gel. These were cloned into a donor vector pDONR201 (Invitrogen, USA) and then transferred into a baculovirus transfer vector pDEST8 (Invitrogen, USA). The recombinant pDEST8 was purified and used to transform DH10Bac-competent cells (Invitrogen, USA), producing recombinant bacmids containing the VP1 gene.

Expression in insect cells

Recombinant bacmids were transfected into Sf9 cells (Riken Cell Bank, Japan) and the recombinant baculoviruses were collected as previously described [10]. The recombinant baculoviruses were used to infect approximately 3×10^6 confluent Tn5 cells (Invitrogen, USA) at a multiplicity of infection (MOI) of 5–10 in 1.5 ml of Ex-Cell 405 medium (JRH Biosciences, USA), and the infected cells were incubated at 26 °C. The culture medium was harvested 5–6 days post-infection (dpi), centrifuged for 10 min at $3,000 \times g$, and further centrifuged for 30 min at $10,000 \times g$. The VLPs were concentrated by ultracentrifugation for 2 h at 45,000 rpm at 4 °C (Beckman TLA-55 rotor), and then resuspended in 30 μ l of Grace's medium. Samples were examined for VLP formation by electron microscopy (EM).

EM

The harvested culture medium was examined for VLPs by negative-stain EM. Briefly, the samples (diluted 1:10 in distilled water) were applied to a carbon-coated 300-mesh EM grid and stained with 4% uranyl acetate (pH 4). Grids were examined in an electron microscope (JEM-1220; JEOL, Japan) operating at 80 kV. VLP images were of CsCl purified culture medium as described previously [10].

Cloning of viral cDNA to produce the 5' and 3' halves of VP1 and ORF2

The Mc114, C12, and NK24 N- and C-terminal regions of VP1 were constructed in order to raise antibodies for the cross-reactivity study. Mc114 recombinant ORF2 (rVP2) was constructed in a similar manner. The primer sequences used to amplify these regions are listed in Table 1. Briefly, the PCR-amplified fragments (using N-terminal sense and antisense

primers, C-terminal sense and antisense primers, or ORF2 sense and ORF2 antisense primers) were cloned into vector pDONOR201 (Invitrogen, USA), and then transferred to vector pDEST17 (Invitrogen, USA) according to the manufacturer's protocol.

Expression in Escherichia coli (E. coli)

pDEST17 plasmids containing N- and C-terminal VP1 and VP2 were transformed into BL21-AI (Invitrogen, USA). Expression was induced by adding 0.2% (w/v) arabinose, followed by incubation at 37 °C for 3 h. The cells were centrifuged for 10 min at $10,000 \times g$ at 4 °C, and were resuspended in extraction buffer (BD Clontech, USA) supplemented with 8 M urea. The supernatant was separated from the cell suspension, and the His₆-tagged recombinant protein was purified in TALON resin (BD Clontech, USA) and finally eluted in buffer containing 250 mM imidazole (BD Clontech, USA). The quantity of protein was estimated using the Protein Assay Kit (BioRad Laboratories, USA).

Time-course expression of Mc114 in insect cells

The expression of Mc114 in the culture medium was analyzed by infecting Mc114 recombinant baculoviruses at a MOI of 14.5 in 2.7×10^6 confluent Tn5 cells in 1.5 ml of Ex-Cell 405 medium followed by incubation at 26 °C. The culture medium was harvested 1, 2, 3, 4, 5, 6, 7, and 8 dpi, centrifuged for 10 min at $3,000 \times g$, and further centrifuged for 30 min at $10,000 \times g$. The VLPs were concentrated by ultracentrifugation for 2 h at 45,000 rpm at 4 °C (Beckman TLA-55 rotor) and resuspended in 20 μ l of Grace's medium. Western blotting, EM, and an antigen ELISA were used to monitor the expression levels.

SDS-PAGE

We examined the rVP1 and rVP2 expression using SDS-PAGE with a 5–20% gradient polyacrylamide gel (ATTO, Japan). The concentrated culture medium and cell lysate (diluted 1:10 in distilled water) were mixed with a 1/4 volume of buffer solution containing 62.5 mM Tris-HCl (pH 6.8), 25% (w/v) glycerol, 2% (w/v) SDS, and 0.01% Bromophenol Blue with 5% (v/v) 2-mercaptoethanol and then boiled for 5 min. Electrophoresis was performed in 25 mM Tris/192 mM glycine/0.1% SDS buffer at 20 mA for 1.5 h.

Western blotting

The proteins were separated by SDS-PAGE and electrotransferred to PVDF with transfer buffer (25 mM Tris/192 mM glycine/5% methanol) at 100 mA for 1 h and blocked with 5% (w/v) skim milk/PBS for 1 h. Proteins were detected with hyperimmune rabbit antiserum at a dilution of 1:3,000 (as determined previously), then following the manufacturer's instructions were developed by chemiluminescence using ECL detection reagent (Amersham Biosciences, England).

Antibody production

Rabbits and guinea pigs were immunized subcutaneously with 10 μ g of CsCl-purified Mc114 VLPs as described elsewhere [10]. For *E. coli*-expressed proteins, 500 μ g of each recombinant protein was used to immunize rabbits at two-week intervals. The serum was collected one week after the last injection.

ELISA

The wells of 96-well microtiter plates (Maxisorp, Nunc, Denmark) were each coated with 100 μ l of a 1:8,000 dilution (determined previously) of either Mc114 VLP hyperimmune

rabbit antiserum (P) or preimmune rabbit antiserum (N) diluted in PBS. The plates were incubated overnight at 4 °C. The wells were washed three times with PBS containing 0.1% Tween 20 (PBS-T) and then were blocked with PBS containing 5% skim milk (PBS-SM) for 1 h at room temperature. The wells were washed four times with PBS-T. Five micro-liters of VLPs (see expression in insect cells section) were mixed in 400 μ l of PBS-T containing 1% SM (PBS-T-SM), and then 100 μ l of this mixture was added to duplicate wells. The plates were then incubated for 1 h at 37 °C. After the wells were washed four times with PBS-T, 100 μ l of a 1:8,000 dilution of Mc114 VLP hyperimmune guinea pig antiserum diluted in PBS-T-SM was added to each well, and the plates were incubated for 1 h at 37 °C. The wells were washed four times with PBS-T, and then 100 μ l of a 1:1,000 dilution of horseradish peroxidase (HRPO)-conjugated rabbit anti-guinea pig immunoglobulin G (IgG) diluted in PBS-T-SM was added to each well. The plates were then incubated for 1 h at 37 °C. The wells were washed four times with PBS-T, and then 100 μ l of substrate *o*-phenylenediamine and H₂O₂ was added to each well and left in the dark for 30 min at room temperature. The reaction was stopped by the addition of 50 μ l of 2 M H₂SO₄ to each well, and the absorbance was measured at 492 nm (A_{492}). For this experiment, we included Tn5 cell lysate and native baculovirus as negative controls. We determined the mean P/N ratio of ELISA was 0.94, with a standard deviation of 0.09 (manuscript in review). The cutoff value of ELISA was defined as the mean plus 3 standard deviations, hence a sample with an A_{492} (P–N) of >0.1 and a P/N ratio of >1.2 was considered significantly positive. For the antigen ELISA, CsCl-purified Mc114 VLPs were used as the positive control at concentrations ranging from 500 to 0.24 ng (data not shown).

Results

Genetic analysis

The nucleotide sequence of the 3' end of the genome (containing the VP1 gene) was determined for each of the strains, Mc114, C12, and NK24. Based on the recent SaV classification [6], these strains belonged respectively to the genogroups GI, GII, and GV (Fig. 1). The Mc114 VP1 encoded 561 amino acids and had an apparent molecular weight of approximately 60,100 (60K protein); the C12 VP1 encoded 561 amino acids and had an apparent molecular weight of approximately 60,100 (60K protein); and NK24 VP1 encoded 569 amino acids and had an apparent molecular weight of approximately 60,500 (60K protein). Figure 2 shows the VP1 amino acid alignments of these strains. All of the sequences contained the predicted VP1 start amino acid motif, MEG (Fig. 2). Based on the recent structural analysis of SaV GI VLPs [4], the Mc114, C12, and NK24 VP1 amino acids were predicted to be more conserved between the 5' to P2 and P2 to 3' domains than in the P2 domain (Table 2). Mc114 and NK24 ORF2 (VP2) encoded 161 and 155 amino acids, respectively, and shared 41% amino acid similarity.

EM analysis

The insect cell culture medium was harvested at 5–6 dpi and examined for VLPs by negative-stain EM. Mc114, C12, and NK24 rVP1 all formed VLPs with diameters of 41–48 nm, while NK24 rVP1 also formed smaller VLPs with diameters of 26–31 nm, though these made up a smaller proportion than the 41–48 nm VLPs (Fig. 3). The 41–48 nm VLPs were morphologically similar to native SaV,

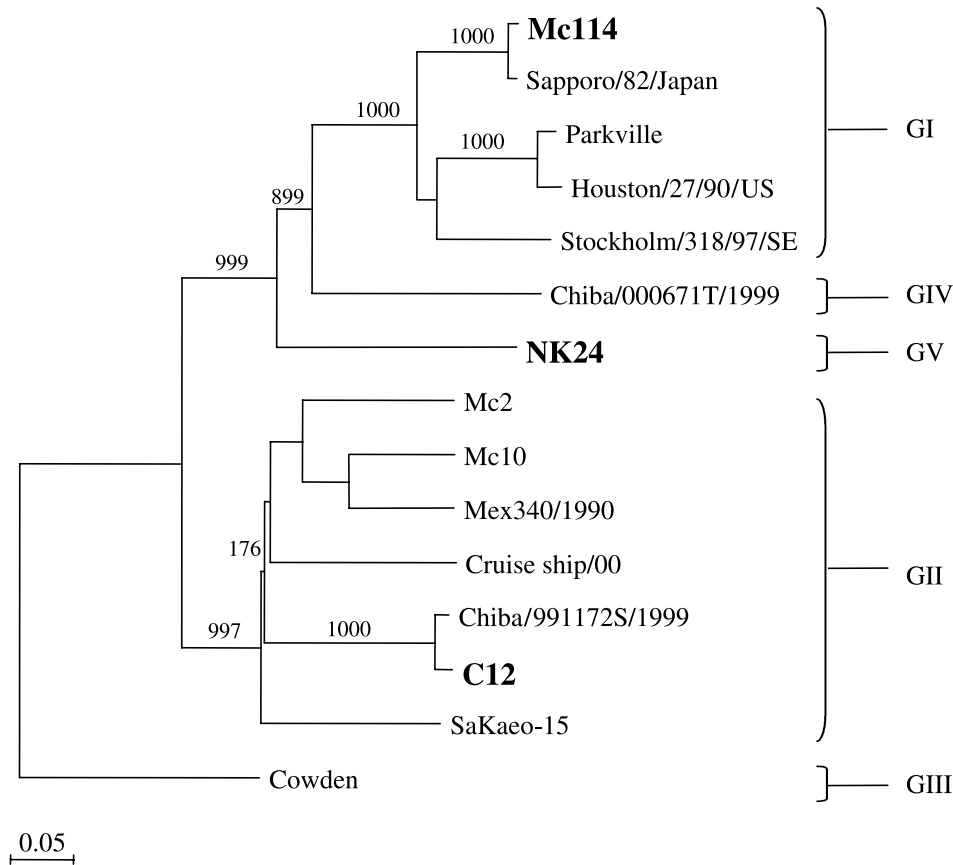


Fig. 1. Phylogenetic analysis of Mc114, C12, and NK24 strains (bold) based on capsid sequences (376 bp). SaV were classified based on the scheme of Farkas et al. [6], using Cowden (SaV GIII) as an outgroup. The number on each branch indicates the bootstrap value for the genotype. Bootstrap values of 950 or higher were considered statistically significant for the grouping [15]. The scale represents nucleotide substitutions per site. GenBank accession numbers for the reference strains are as follows: Chiba/000671T/1999, AJ412805; Chiba/991172S/1999, AJ412797; Cruise ship/00, AY289804; Houston/27/90/US, U95644; Mc2, AY237419; Mc10, AY237420; Mex340/1990, AF435812; Cowden, AF182760; Parkville, U73124; Sakaao-15, AY646855; Sapporo/82/Japan, U65427; and Stockholm/318/97/SE, AF194182

including the typical Star of David structure, cup-shaped surface depressions, and ten spikes on the outline. However, the Star of David structure was visible only when the samples were stained with uranyl acetate, and not with phosphotungstic acid (data not shown). The 26–31 nm VLPs had spikes on the outline, but neither the Star of David structure nor cup-shaped surface depressions were visible.

Time-course expression analysis of Mc114 rVP1 in insect cells

Three tests were used to monitor the time-course expression of Mc114 rVP1 in insect cells. The antigen ELISA first detected VLPs in the culture medium at 3 dpi (Fig. 4A). The Western blot with hyperimmune rabbit Mc114 VLP antiserum first detected rVP1 (60K band) at 4 dpi (Fig. 4B). And VLPs were first detected by EM at

Mc114 1 MEGNGSNSEPKQSNPNMV----VDP---PGTT-GPTTSHVVVANPEQPNGAAQRLEAVATGAIQSNVPEAIRNCFAVFRFTFAWDRMPTGTFLGSLH 92
NK24 1RLGNS.TQSH.DTSNT.QGAP-....-.ADAPL.PV.....LP.....I...TS...DCV.S...LL..IP..T.Q.Q.SL.TAV... 98
C12 1 ..-----VPRPEG.KANSNENV.LAS.QD.I..NAALLPTQI.T.....V.M.A...VSN...MCV.E...SVT.LP.TT.QASN...A.H.G 94
*** *

Mc114 93 PNINPYTSHLSGMWAGWGGSEVRLSISGSGVFAGRIIASVIPGVDPSIRDGPVLPFAFVDARITEPVFSMIPDVRAVDYHRMDGAEPTCSLGFVWVYQ 192
NK24 99 .D....K..AQ.F...AMDI.VTV...L...KLVCG.L...N.TLVN.....L.....AC.NVA.....T..D.A.AT..I..L 198
C12 95 .R....A...A.F.....QI.VTL...LY...AVVA.L...N.ANVQN...F...I...TVD.ILINL.I.....V..D.Q.ATV.L..A 194
* ***** ** * ***** * ***** ** * ***** * ***** ** * ***** ** * ***** ** * ***** *

Mc114 193 PLLNPFSTTAVSTCWVSVETKPGGDFDFCLLRPPGQMQENGVSPEGLLRRLGYSRGNVGGVVGGMILVAEHKQVNRHFNSSVTF**GWSTAPVNPMAAE** 292
NK24 199 ..I....NE...A.I.I.....L...K...A...G..SH...K.QRA...A..YA...VI.GSAH.....TALGT.....YE..RCA 298
C12 195 .I...Q.GSI...LTF..R.P.....KA.E.E.D..I..AN.....R....L.R...LVV..VAE...H..AA..T.L...L.IE.I.GA 294
** ** *

Mc114 293 **I**V**T**NQ**A**H**S**T**S**R**H**--**A**--**W**L**S**I**G**A**Q**N**K**G**L**P**G**I**P**N**H**F**P**D**S**C**A**S**T**V**G**A**M**D**T**S**L**G**R**P**S**T**G**V**C**----**G**P**A**I**S**F**Q**N**G**D**V**Y**E**N**D**T**P**S**V**M**F**A**T**-**Y**D**P**L**T**S**G**-**T** 382
NK24 299 FGGVHVA.G.TPKKIYGY.E-V..DQR....N.V..W..FAVNSKYSWP.ADYI--.YNA.V----.TLV..AD....S..EVATA-..VSMNPST..GR 390
C12 295 .SWY.NTPPGIST----RGLLS.EG..II..N.V..WT.VAL.SKTSGR.T.VPTDQANLNQ.PGAS..VVM.....N.TSANNVCLTAASHDFVNLS 390
* *

Mc114 383 **G**V**A**L**T**N**S**I**N**P**A**S**L**A**L**V**R**I**S**N**N**----**D**F**D**T**S**G**F**A**N**D**K**N**V**V**Q**M**S**W**E**M**Y**T**G**T**N**Q**I**R**G**Q**V**T**P**M**S**G**T**I**N**Y**T**F**T**S**T**G**A**N**T**L**V**L**W**Q**E**R**M**L**S**Y**D**G**H**Q**A**I**L**Y**S**S**Q**L**E**R**T** 478
NK24 391 .T..RENFD.STMH...TNSTAQPA.WPS-.SNTNGYFTP,-.GHGS.NA-.NDK..N.E.A...GGS.Q.NI...I.KIF.DHPGPT.....DS. 486
C12 391 NFDAAAGMWW----.PWTTTK--PDATINRNVYITPT-----INGDPSRP.H.KC.N.V..FQ.GG..T.NIM...QHFTSFPGA.EV.C....S. 478
* *

Mc114 479 AEYFQNDIVNIPENSMAVFNVTNSASFQIGIRPDGYMVTGGSIGINVPLEPETRFQYVIGILPLSALS GP SGPNMGRA-----KRVFQ 561
NK24 487 .TI..SGP.....M...Y..T..G.D..V...R.....S.T..TRQE.D.D.T.T...LF...S.V..H..T...--QIAWS--- 569
C12 479 .M...NV...A.Q.....AGNT...A.M.N..C..NAAV.THQL.DY..S.RF..LF.Q.TS.Q..N..A...VRFLE----- 561
* *

Fig. 2. An amino acid alignment of VP1 sequences of Mc114 (GI), C12 (GII), and NK24 (GV) strains. The sequences with a rectangular box represent the predicted P2 domain [4]. The asterisks indicate conserved amino acids among these three VP1 sequences

Table 2. VP1^a amino acid (%) identities of SaV Mc114, C12, and NK24

	Mc114				C12			
	VP1	5' to P2	P2	P2 to 3'	VP1	5' to P2	P2	P2 to 3'
C12	46.0	55.0	26.7	51.6				
NK24	51.0	60.0	29.9	59.0	50.1	59.8	31.8	55.0

^aDomains and regions were based on Fig. 2

4 dpi. The expression level of VLPs increased each day thereafter, peaking at 6 dpi before decreasing at 7 and 8 dpi (Fig. 4A). A thin band of approximately 55K was also detected by Western blotting at 4 dpi, which increased each day thereafter as evidenced by an increase in band intensity (Fig. 4B). The 55K band may have represented truncated form(s) of rVP1 as observed with norovirus expression studies [2], though direct evidence is lacking. All bands were absent in the cell controls (the wild-type baculovirus and the mock-infected Tn5 cells) (data not

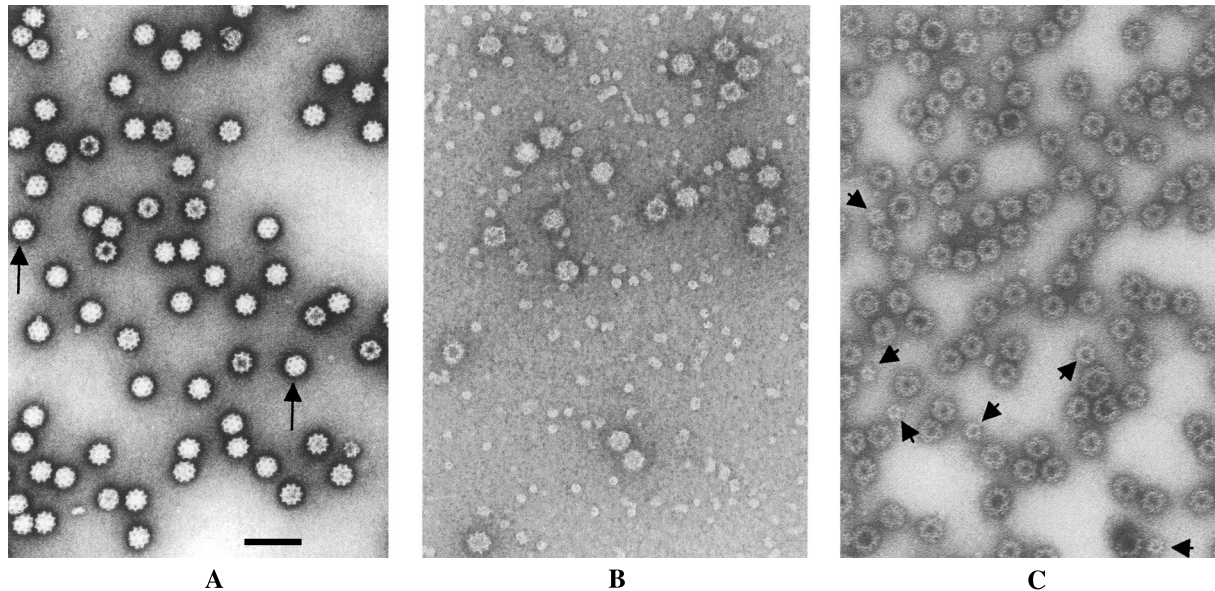


Fig. 3. Electron-microscopic images of CsCl purified (A) Mc114, (B) C12, and (C) NK24 VLPs negative-stained with 4% uranyl acetate (pH 4). The long arrows show the Star of David structure and the short arrows show the small VLPs. The bar indicates 100 nm

shown). Our results indicated that the optimal time to harvest Mc114 VLPs in culture medium was 6 dpi (Fig. 4A).

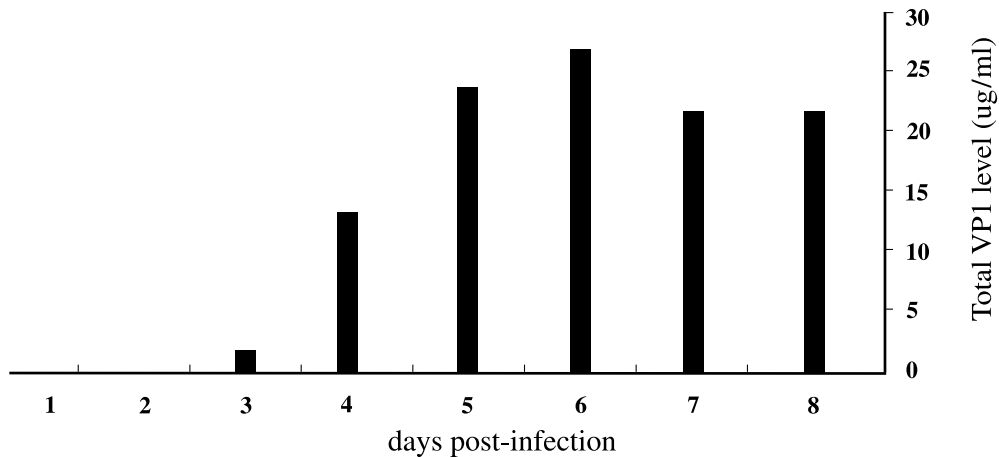
We also analyzed the time-course expression of Mc114 rVP2 in insect cells by Western blotting with antiserum raised against *E. coli*-expressed Mc114 VP2. However, no bands were detected in either the culture medium or the cell lysate (data not shown).

Western blot cross-reactivity analysis

Western blotting was used to investigate the cross-reactivities among SaV Mc114, C12, and NK24 rVP1 expressed in insect cells. Hyperimmune rabbit antiserum was raised against *E. coli*-expressed Mc114, C12, and NK24 N- and C-terminal VP1. In addition, hyperimmune rabbit antiserum raised against Mc114 VLPs was used. As shown in Fig. 5 and summarized in Table 3, Mc114 (VLP, N- and C-terminal) antisera were reactive with Mc114 and NK24 rVP1 (Mc114 VLP antiserum weakly with NK24 rVP1) but not with C12 rVP1. On the other hand, C12 (N- and C-terminal) antisera were reactive with C12 rVP1 and weakly reactive with Mc114 rVP1, but were not reactive with NK24 rVP1, whereas NK24 (N- and C-terminal) antisera reacted with NK24 rVP1 only.

The three kinds of Mc114 antisera reacted in manner similar to that of Mc114 rVP1, revealing bands at 60K and 55K (Fig. 5), as observed in the time-course analysis (Fig. 4B). These two bands were also detected by C12 N-terminal antiserum with Mc114 rVP1 but not with the C12 C-terminal antiserum. One band was detected by the C12 N-terminal antiserum with C12 rVP1, whereas two bands of

A



B

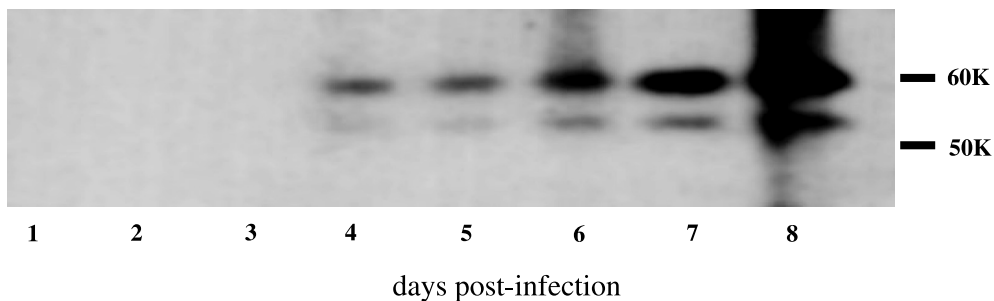


Fig. 4. Time-course expression of the Mc114 rVP1. Confluent Tn5 cells were infected with Mc114 recombinant baculoviruses at MOI of 14.5 and incubated at 26 °C. The culture medium was harvested at the indicated day (days post-infection) and concentrated as described in the Materials and methods. Using the same samples, the expressed proteins were analyzed by (A) antigen ELISA with hyperimmune rabbit (capture) and guinea pig (detector) antisera raised against Mc114 VLPs and (B) Western blotting with hyperimmune rabbit antiserum raised against Mc114 VLPs. For the antigen ELISA, purified Mc114 VLPs were used as the positive control at concentrations ranging from 500 ng to 0.24 ng

approximately 60K and 58K were detected by the C12 C-terminal antiserum. Two bands of approximately 60K and 55K were detected by NK24 (N- and C-terminal) antisera with NK24 rVP1, and an additional weak band of approximately 45K was detected by the NK24 C-terminal antiserum. Two bands of approximately 60K and 55K were detected by Mc114 VLP and N-terminal antisera with NK24 rVP1, but only a single band of 60K was detected by Mc114 C-terminal antiserum. The significance of these lower bands has not yet been determined.

ELISA cross-reactivity analysis

An antigen ELISA was used to examine the cross-reactivities among Mc114, C12, and NK24 VLPs expressed in insect cells. The ELISA incorporated hyperimmune rabbit (capture) antiserum and guinea pig (detector) antiserum raised against purified Mc114 VLPs. Samples were added to duplicate wells and averaged.

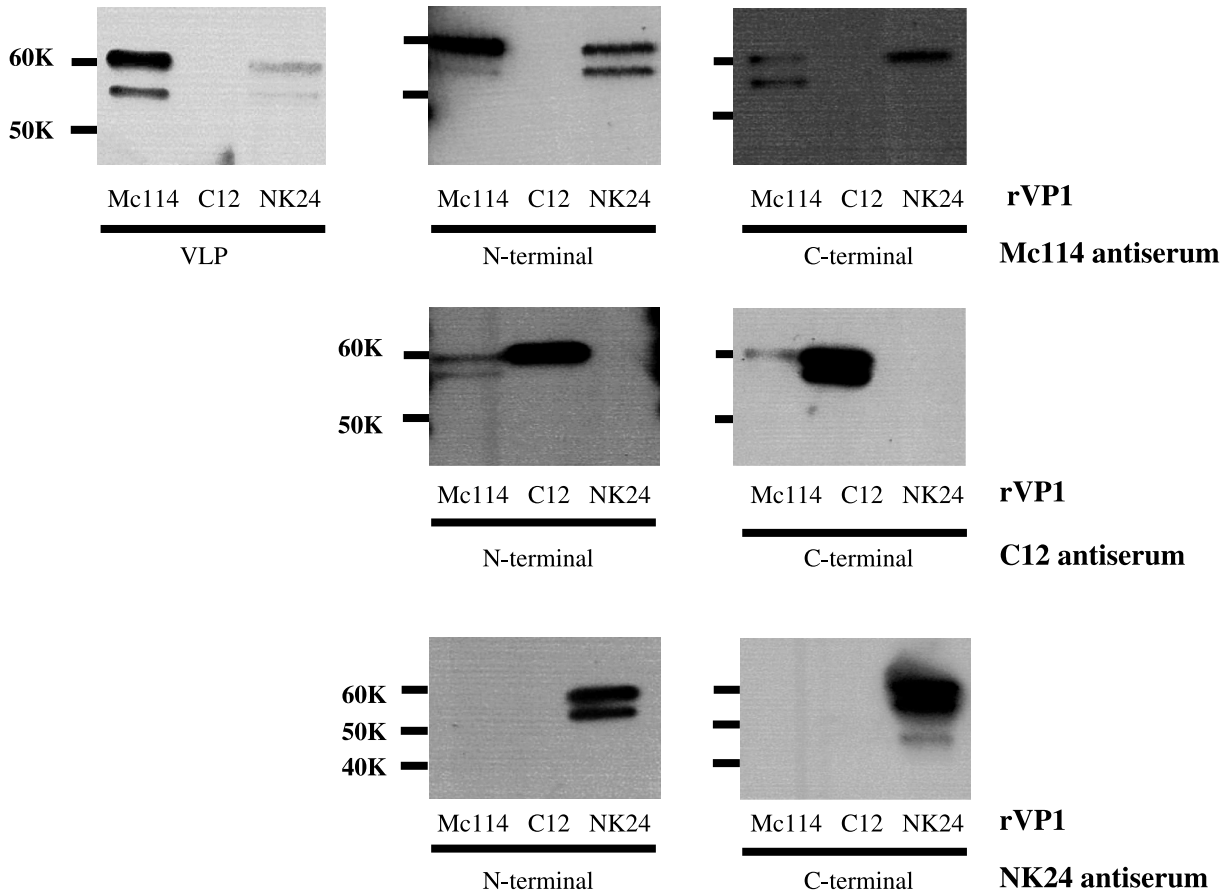


Fig. 5. Western blots of Mc114, C12, and NK24 rVP1 expression in insect cells. Culture medium was harvested at 6dpi. Seven different hyperimmune rabbit antisera were used, including antisera raised against *E. coli*-expressed Mc114, C12, and NK24 N- and C-terminal VP1, and antisera raised against Mc114 VLPs. The band intensities are relative to each antiserum

Table 3. Reactivities of antiserum with SaV GI, GII, and GV rVP1

rVP1	ELISA	Western blot						
	Mc114 antisera ^a	Mc114 antiserum			C12 antiserum		NK24 antiserum	
	VLP	VLP	N-terminal	C-terminal	N-terminal	C-terminal	N-terminal	C-terminal
Mc114 (GI)	0.46 (9.23) ^b	+	+	+	+ ^c	+ ^c	-	-
C12 (GII)	0.00	-	-	-	+	+	-	-
NK24 (GV)	0.00	+ ^c	+	+	-	-	+	+

^aThe ELISA uses Mc114 VLP (rabbit) antiserum as capture and Mc114 VLP (guinea pig) antiserum as detector

^bP-N and (P/N ratio) values measured at 492 nm

^cWeakly reactive

As shown in Table 3, the ELISA detected Mc114 VLPs (A_{492} P-N = 0.46 and P/N ratio = 9.23) but neither C12 VLPs nor NK24 VLPs (A_{492} P-N = 0.00).

Discussion

Human SaV VP1 start contains an amino acid motif, MEG, which is conserved in all human SaV strains and as such is considered to be the putative VP1 start. In this study, we re-examined an earlier suggestion that an upstream sequence from the VP1 start AUG codon is crucial to the formation of human SaV VLP [14]. In addition, we examined the cross-reactivities among SaV GI, GII, and GV rVP1 by Western blotting and the cross-reactivities among SaV GI, GII, and GV VLPs by antigen ELISA. Our results show the following: (i) SaV GI, GII, and GV rVP1 constructs beginning exactly from the predicted VP1 start AUG codon self-assembled VLPs; (ii) an upstream sequence of the predicted VP1 start AUG codon was not an essential element for SaV GI, GII, or GV VLP formation; (iii) the morphological features of the predominant SaV GI, GII, and GV VLPs were similar to those of native SaV, except that GV VP1 also expressed small VLPs; (iv) by Western blot, GI antisera cross-reacted with GV rVP1 but not with GII rVP1, whereas GII antisera cross-reacted weakly with GI rVP1 but did not cross-react with GV rVP1, while GV antisera reacted with GV rVP1 only; and (v) by antigen ELISA, GI VLPs were antigenically distinct from GII and GV VLPs.

Only three other expression studies of human SaV VLP formation have been reported [4, 14, 22]. In one of those studies, an upstream sequence of 73 nucleotides from the predicted VP1 start AUG codon (construct pHou/90-3, Houston/27/90/US strain) was a crucial element for VLP formation [14], whereas one of the other two reports included 39 nucleotides upstream (Sapporo/82/Japan strain) [22]. Recently, the Parkville strain (U73124) was reported to form empty VLPs with a construct that began exactly from the predicted VP1 start. Surprisingly, the Parkville strain had 97.5% and 100% nucleotide identity to pHou/90-3 VP1 and the 73-nucleotide-upstream sequences, respectively. On the other hand, Mc114 had 94.4% and 94.9% nucleotide identity to the Sapporo/82/Japan VP1 and the 39-nucleotide-upstream sequences, respectively. These results suggest that an upstream sequence may not be an essential element for SaV VLP formation, but that some other factor(s) are necessary.

Porcine enteric calicivirus (PEC) of SaV GIII was also expressed using a baculovirus expression system that resulted in the formation of PEC VLPs [9]. The PEC rVP1 construct contained nine nucleotides upstream of the VP1 start AUG codon [9]. That study also included a bacteriophage T7 RNA polymerase promoter of 19 nucleotides (in front of these 9 nucleotides) and a downstream sequence of 98 nucleotides at the 3' of the VP1 gene. Although the T7 promoter is independent of the baculovirus expression system, Jiang suggested that an element (i.e., an upstream sequence) could be responsible for initiating transcription or translation; alternatively, this element could serve as a spacer between the polyhedrin promoter and the VP1 gene, or as a stabilizer of the RNA after transcription [26]. Interestingly, Jiang also tried to express two other recombinant constructs (pHou/90-1

and pHou/90-2) that included 9 and 29 nucleotides upstream, respectively, but those constructs failed to express rVP1 or form VLPs [14].

Human norovirus (NoV) also belongs to the family *Caliciviridae* and is typically associated with food- and waterborne infections [1]. Human SaV and NoV strains are morphologically, genetically, and antigenically distinct [1, 16]. The NoV genome is organized in a slightly different way than the SaV, since ORF1 encodes all the nonstructural proteins, ORF2 encodes the capsid protein (VP1), and ORF3 encodes a small protein (VP2). In a recent NoV expression study, inclusion of VP2 and poly(A) sequences in NoV rVP1 constructs were found to stabilize VLP formation [3]. In the pHou/90 constructs, both VP2 and poly(A) sequences were absent [14]. Consequently, if SaV VP2 has functions similar to those of NoV VP2, as has been suggested [1], then some crucial factors for SaV VLP formation could be SaV VP2 and poly(A) sequences. Although in a recent report the rVP1 was expressed for a SaV GII strain as determined by SDS-PAGE [24], VLPs were not formed even when an upstream sequence from VP1 start AUG codon, VP2, and poly(A) sequences were included in the construct.

All of our rVP1 constructs formed the typical native SaV Star of David structure and had diameters of 41–48 nm, while NK24 rVP1 also formed smaller VLPs with diameters of 26–31 nm (Fig. 3). The recombinant Sapporo/82/Japan construct had the Star of David form [22], whereas the pHou/90-3 construct did not [14]. Besides the different genogroups and genotypes, the only obvious differences in the constructs that formed this structure were the inclusions of VP2 and poly(A) sequences. These results suggest that VP2 and/or poly(A) sequences are needed to form the Star of David structure, although direct evidence is lacking.

The Mc114 rVP2 was not detected by our Western blotting during the time-course analysis (data not shown), although NoV studies have found that the expression level of NoV rVP2 was low [3]. On the other hand, we did not detect any SaV rVP1 cleavage products either (data not shown), and a similar result was observed in other SaV expression studies [9, 14, 22], whereas for NoV the rVP1 is cleaved at amino acid residue 227, yielding a 32K C-terminal rVP1 product [12, 16]. These data suggest that SaV rVP1 expression could be quite different from that of NoV. Our studies have also found that the yields of purified SaV VLPs were low in comparison with those of NoV VLPs (data not shown), which was also discussed by Guo et al. [9].

Hyperimmune rabbit and guinea pig antisera raised against GI VLPs were used to examine the cross-reactivities among SaV GI, GII, and GV VLPs by an antigen ELISA. As summarized in Table 3, the antigen ELISA was specific only for the homologous GI VLPs. This result was not so unusual, since the amino acid homologies among the VP1 sequences were low (Fig. 2). In the recent structural analysis of SaV VLPs, the outermost domain of VP1, i.e., the P2 domain, was shown to be the most variable region and is likely to confer strain diversity [4]. The predicted P2 domain was the most variable region among these three SaV VP1 sequences, whereas the S and P1 domains were more conserved (Table 3). These data suggest that there were no shared epitopes among GI, GII, and GV VLPs as determined by antigen ELISA, though further studies are needed to confirm this suggestion.

Also, hyperimmune rabbit antisera were raised against *E. coli*-expressed GI, GII, and GV N- and C-terminal VP1 in order to examine the cross-reactivities among SaV GI, GII, and GV rVP1 by Western blotting (Fig. 5 and Table 2). We found that the GI antisera cross-reacted with GV rVP1, but that GV antisera did not cross-react with GI rVP1, i.e., there was only a one-way cross-reactivity. Also, GII antisera cross-reacted weakly with GI rVP1, but GI antisera did not cross-react with GII rVP1, i.e., this too was a one-way cross-reactivity. Figure 2 shows that short continuous residues (8–9 amino acids) had 100% homology between VP1 sequences. These short residues may have represented particular target residues for the polyclonal antibodies that were raised against *E. coli*-expressed VP1. A similar result was observed by Yoda et al. [28] with *E. coli*-expressed NoV VP1. Yoda suggested specific conformational epitope(s) or limited continuous epitope(s) in the NoV VP1 residues that allowed for broad cross-reactivity between NoV GI and GII VP1, which generally have a low amino acid homology. The reason(s) for these SaV one-way cross-reactivities have not yet been determined, nor has the significance of the double (and triple) bands by Western blotting (Fig. 5). Comparable double bands are usually detected for NoV rVP1 expression and were found to include N-terminal cleavage products [2, 16]. However, these SaV extra bands have not yet been determined.

Interestingly, GI VLP rabbit antiserum cross-reacted weakly with GV rVP1 but did not cross-react with GII rVP1 by Western blotting (Fig. 5). Farkas et al. [6] showed that the SaV GV VP1 sequence was related more to GI than GII, based on phylogenetic distance analysis. Also, GI and GV strains possess a predicted ORF3 (VP3?), whereas the ORF3 is absent in GII strains. Further investigations are clearly needed to determine the significance of these novel findings.

Acknowledgments

This work was supported by grants-in-aid from the Ministry of Education, Culture, Sports, Science and Technology of Japan and by a grant for Research on Re-emerging Infectious Diseases from the Ministry of Health, Labour, and Welfare of Japan. We are grateful to the Japanese Monbusho for the PhD scholarship provided to G. Hansman. We thank Mr Hatano for his assistance with the EM.

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