Molecular Characterization of Infectious Clones of the Minute Virus of Canines Reveals Unique Features of Bocaviruses

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Minute virus of canines (MVC) is a member of the genus Bovavirus in the family Parvoviridae. We have molecularly cloned and sequenced the 5′- and 3′-terminal palindromes of MVC. The MVC genome, 5,404 nucleotides (nt) in length, shared an identity of 52.6% and 52.1% with that of human bocavirus and bovine parvovirus, respectively. It had distinct palindromic hairpins of 183 nt and 198 nt at the left-end and right-end termini of the genome, respectively. The left-end terminus was also found in two alternative orientations (flop or flop). Both termini shared extensive similarities with those of bovine parvovirus. Four full-length molecular clones constructed with different orientations of the left-end terminus proved to be infectious in Walter Reed canine cell/3873D (WRD) canine cells. Both MVC infection and transfection of the infectious clone in WRD cells revealed an identical RNA transcription profile that was similar to that of bovine parvovirus. Mutagenesis of the infectious clone demonstrated that the middle open reading frame encodes the NP1 protein. This protein, unique to the genus Bovavirus, was essential for MVC DNA replication. Moreover, the phospholipase A2 motif in the VP1 unique region was also critical for MVC infection. Thus, our studies revealed important information about the genus Bovavirus that may eventually help us to clone the human bocavirus and study its pathogenesis.

Bovavirus is a newly established genus in the subfamily Parvovirinae of the Parvoviridae family (61). It consists of two formal members, bovine parvovirus (BPV) and minute virus of canines (MVC) (61), and one tentative member, the human bocavirus (HBoV) (4). MVC was first recovered from canine fecal samples in 1970 (8). Although it was initially thought not to cause disease, recent studies of experimental infection revealed pathogenicity for newborn pups and fetuses (10). It caused enteritis with severe diarrhea (9, 44) and respiratory disease with breathing difficulty (31, 52); however, these symptoms were often associated with coinfection of other viruses (44). Pathological lesions in fetuses in experimental infections were found in the lungs and small intestine (10). Pathology caused by MVC was seen most commonly in animals between the ages of 1 to 5 weeks (10, 29, 31, 52). Serological evidence indicated that MVC was widespread in the dog population, with a 50 to 70% seroprevalence rate worldwide (11).

A virus that is associated with lower respiratory tract infections in humans was identified in 2005 and named HBoV because of its high similarity to BPV and MVC in its genomic sequence (4). The HBoV genome was detected in 1.5% to 19% of respiratory specimens from symptomatic hospitalized children (3, 7, 39). Most children infected with HBoV were younger than 24 months (3, 7, 39). HBoV was also detected against HBoV capsid suggested that children were infected with HBoV as early as 3 months of age, and most infections occurred before 1 year of age (25, 33, 34, 38). Isolation and culture of the virus have not been reported so far.

Our current knowledge of the genus Bovavirus is obtained mostly from studies of BPV. The BPV genome was sequenced some 20 years ago and is composed of 5,515 nucleotides (nt) (12, 13, 54). Three capsid proteins, VP1 (80 kDa), VP2 (72 kDa), and VP3 (62 kDa), have been detected in purified virions as well as during viral infection (32, 35). Analysis of the genomic sequence (13) revealed three open reading frames (ORFs). The left-hand ORF was predicted to encode the non-structural protein NS1. The middle ORF (mid-ORF) of BPV is thought to encode the abundant 24-kDa nonstructural protein NP1, whose function is largely unknown (36). The right-hand ORF contains the coding sequences for the overlapping capsid protein genes VP1 and VP2. Only one promoter at a map unit of 4 (P4) was identified that transcribes one pre-mRNA. This pre-mRNA is processed to generate at least eight transcripts through alternative splicing as well as alternative polyadenylation (54).

MVC was isolated and grown in the Walter Reed canine cell/3873D (WRD) cell line (8). An incomplete sequence of the genome of MVC that lacked critical sequences of the palindromic repeats (approximately 150 nt at both ends) was previously obtained (GenBank accession no. NC_004442) (58). This sequence showed 43% identity to that of BPV, while the NS1, VP1, and NP1 proteins were 33.6%, 41.4%, and 39% identical to those of BPV (58). The sequence of another MVC isolate (HM-6; GenBank accession no. AB158475) that also lacked both the left and right termini showed 96.3% identity to that of the prototype MVC sequence (48). An infectious clone for mutational studies is necessary to fully understand the molecular aspects of the genus Bovavirus and especially viral...
DNA replication. Although an infectious clone of BPV was reported (59), it was no longer available. Unfortunately, the palindromic repeats of HBoV are currently unknown. This seems to be due to difficulties in obtaining large quantities of virus, either by growing the virus in tissue culture or from clinical specimens that contain large quantities of virus, which are necessary to clone them. Because of similarities between HBoV and MVC both in the organization and sequence of the genomes and in the symptoms of diseases they cause, we decided to clone the MVC palindromic termini and thereafter construct the infectious clones described in this report. Characterizing a selected infectious clone provided us invaluable information about DNA replication and the functions of NP1 and VP1 of baculoviruses in general.

MATERIALS AND METHODS

Virus and cells. The MVC virus used in this study was the original strain (GA3) isolated from lung tissue (10). It was grown in the WRD cell line (8) through two passages (MVC GA3 P2). Both the virus and the cell line were gifts from Colin Parrish at the James A. Baker Institute of Cornell University. The virus-containing supernatant was adjusted to a density of approximately 1.40 g/ml contained the highest genomic copy number per ml and was used to prepare dsDNA of BPV as described above. Based on the previously published sequence of the genome (accession no. DQ335247), annealed dsDNA was blunted and digested with EcoRI.

Virus purification and quantification. MVC was used to infect WRD cells in 20 flasks (175 cm²) at a multiplicity of infection of approximately 10³ genomic copies/cell. The infected cells were harvested after the appearance of a clear cytopathic effect and suspended in 10 ml of phosphate-buffered saline. After freezing and thawing three times, released viruses were collected as supernatant with high-speed centrifugation (10,000 rpm for 1 h in a Sorvall RC-6 centrifuge).

Virus-containing supernatant was adjusted to a density of approximately 1.40 g/ml by adding solid CsCl and was spun at 36,000 rpm (Sorvall TH641) for 36 h at 20°C. Fractions were collected in 0.5-ml increments and dialyzed against phosphate-buffered saline with subsequent quantification of MVC genomic copies by a real-time PCR assay. The MVC real-time PCR assay for absolute MVC genomic copies was developed following a previously described method (53) with a TaqMan probe of 5'-6-carboxytetramethylrhodamine-6-carboxyfluorescein-AACACACAAAGCGCGGCTAC TCGG-3' (nt 4376 to 4397 and nt 4401 to 4424), a forward primer of 5'-AGGACCATGCTGTGCTTACT-3' (nt 4376 to 4397), and a reverse primer of 5'-TACTGTCGAGGCTTGGTT-3' (nt 4445 to 4426). The fraction with the density of ~1.39 g/ml contained the highest genomic copies of MVC at 4 × 10⁷ per μl and was used to anneal the single-strand DNA (ssDNA) genome of MVC. All nucleotide numbers in this study refer to the full-length MVC genome (GenBank accession no. FJ214110). Northern blot analyses were done exactly as previously described (50), using MVC RNA as a probe.

Expression plasmids. MVCNP1 was constructed by inserting MVC NP1 ORF (nt 2517 to 3096) plus a C-terminal flag into pCDNA3, while HBoVNP1 was constructed by inserting the HBoV NP1 ORF (nt 2410 to 3067; accession no. DQ000496) plus a C-terminal HA into pcDNA3. The MVC clones were digested with HindIII and EcoRV and were ligated to SpH1 (blunted) HindIII-digested MVC M13 helper plasmids, which resulted in full-length clones of MVC with various right-end and left-end termini in the flip or flop orientation. MVC M13 helper plasmids were deleted in pMVC.

Mutants based on the infectious clone of pMVC. All mutants based on pMVC and the detailed mutations therein are diagrammed (see Fig. 4 and 5, respectively).

P5 RNA probe construct. The P5 probe was cloned by Northern blot analysis of WRD cells. The probe was PCR-amplified from the MVC genome and was inserted into the vector pGEM3Z (Promega, Madison, WI).

Plasmid construction. Plasmids were constructed by inserting a fragment of MVC into BamHI/HindIII-digested vector pGEM4Z (Promega, Madison, WI).

Sequence analyses. Sequence analyses were performed using the BLAST program (www.ncbi.nlm.nih.gov/BLAST).

RESULTS

Cloning and sequencing of the palindromic termini of MVC. A total of 100 μl of MVC at 10³ genomic copies/μl was heated at 95°C for 5 min and cooled slowly to form the double-stranded DNA (dsDNA) of the MVC replication form. In light of the previously published sequence of the MVC NSCap gene (accession no. NC_004442) (58), there are two BspEI sites on MVC genome, one site at nt 210 after the left-end terminus and another one at nt 4469 before the right-end terminus. Thus, annealed dsDNA was blunted and digested with BspEI. BspEI-digested DNA fragments were ligated into the Smal/BspEI-digested vector of pBB1.4A. Positive clones were collected as MVCright clones or MVCleft clones based on sequence results using two sequencing primers that flanked the inserted terminus in pBB1.4A. Preliminary sequencing results revealed that there was a BglI site on the left-end terminus and an AseI site on the right-end terminus as shown in Fig. 1. Next, BglI-digested MVCleft clones were sequenced with the two sequencing primers from both directions, which gave rise to the whole sequence of the left-end terminus. Similarly, AseI-digested MVCright clones were sequenced for the right-end terminus.

The pBB1.4A cloning vector was constructed by inserting a DNA fragment that was composed of a XhoI-EcoRV-NotI-Smal-linked 1.4-kb DNA fragment amplified from the adenosine virus 5 E1A gene (nt 1013 to 2400; accession no. AC_000008) with a right-hand linker of BspEI-SphI-Kpn-HindIII-XhoI into Sall-digested pM20, the B19 infectious clone (68). The forward sequencing primer that was located approximately 176 nt before the insertion described above was 5'-CTGCGCGTAACCACCACA-3'. The reverse sequencing primer that was about 134 nt after the insertion described above was 5'-CATGGGGT CAGTTGGGAC-3'. All clones containing terminal repeats were sequenced by MCLAB (www.mclab.com) using the selected method for the GC-rich/hairpin structure.

(ii) Sequencing the palindromic termini of BPV. A total of 100 μl of purified BPV at 10³ genomic copies/μl was used to prepare dsDNA of BPV as described above. Based on the previously published sequence of the genome (accession no. DQ335247), annealed dsDNA was blunted and digested with EcoRI. There are two EcoRI sites on BPV, one at nt 978 after the left-end terminus and another one at nt 5067 before the right-end terminus. EcoRI-digested DNA fragments were ligated into the SmaI/EcoRI-digested vector of pBB1.4B. Positive clones were collected as BPVRight clones or BPVLeft clones based on sequence results using the two sequencing primers. Preliminary sequencing results showed a BglI site on the left-end terminus and an Ndel site on the right-end terminus as shown in Fig. 1. Finally, the whole sequence of the left-end hairpin was obtained by sequencing Smal-digested fragments of BPVleft clones, and the right-end hairpin sequence was obtained by sequencing Ndel-digested fragments of BPVRight clones.

To construct BPV constructs, clones containing an insert of bpA (nt 321 to 392) were ligated into SmaI/BglI vector pBB1.4A (59). Sequencing results revealed that there was a BglI site on the left-end terminus and an NdeI site on the right-end terminus as shown in Fig. 1. The whole sequence of the left-end hairpin was obtained by sequencing Smal-digested fragments of BPVleft clones, and the right-end hairpin sequence was obtained by sequencing Ndel-digested fragments of BPVRight clones.
A

MVC LEH

\[
\Delta G = -92.22 \text{ kcal/mol}
\]

Flip

Flop

\[
\Delta G = -96.75 \text{ kcal/mol}
\]

BPV LEH

\[
\Delta G = -75.35 \text{ kcal/mol}
\]
cells as described previously (28), and Southern blots were performed as described previously (50) with the MVC NSCap probe diagramed in Fig. 3. Signals of RNase protection assays, Northern blots, and Southern blots were quantified with the Typhoon 941 phosphor imager and ImageQuant TL software (GE Healthcare).

Production of antibody against MVC NS1 and immunofluorescence assay. Glutathione S-transferase (GST)-fused NS1 C-terminal 112 amino acids (GST-NS1c112aa) was expressed in Escherichia coli BL21 cells transformed with pGEXNS1c112 and was purified following instructions from the GST fusion system handbook (GE Healthcare). A GSTrap FF 1-ml column was used for purification of the GST fusion protein by the Biologic LP system (Bio-Rad). The purified GST fusion NS1 protein was analyzed by sodium dodecyl sulfate-polyacrylamide gel electrophoresis (SDS-PAGE) and appeared as a single band at a size of approximately 47 kDa. Purified GST-NS1c112aa (500 μg/ml) was emulsified with TiterMax Gold adjuvant (Norcross, GA) at a ratio of 1:1 and was injected into three Sprague-Dawley rats subcutaneously at a dose of 400 μg/rat.

FIG. 1. Sequences and structures of MVC palindromic repeats. (A) The sequence of the MVC and BPV left-end hairpin (LEH) is shown in flip and flop orientations. The left-end terminus of MVC in flip or flop and the left-end terminus of BPV in flop were terminated at various ends as shown by arrows with an indication of their ended nucleotide numbers. The left-end terminus of BPV shown in flip refers to the previously published sequence (12, 54). The corrected nucleotide in BPV left-end terminus is indicated by the arrowhead. The putative NS1 binding site of MVC is indicated. The rabbit-ear structure that shares a great identity between MVC and BPV is shown. BglI and SmaI that were used to digest the left-end end hairpin for sequencing are shown. (B) Sequences of the right-end hairpin (REH) of MVC and BPV are shown in a stem or stem-arm structure. The right-end terminus of MVC is shown in a stem or stem-arm structure. The right-end terminus of BPV is shown in flip or flop orientation. The stem-arm structure of BPV is shown only in the flop orientation. The right-end terminus was terminated at various ends as shown by arrows with an indication of their respective nucleotide numbers. AseI and NdeI that were used to digest the right-end hairpin for sequencing are shown. The corrected nucleotide in BPV right-end terminus was indicated by the arrowhead. The DNAMAN program (Lynnon, Co., Quebec, Canada) was used to predict the secondary structures of both termini of MVC and BPV.
Six weeks later, blood was taken from the tail vein. Sera were tested for MVC NS1-specific antibody by enzyme-linked immunosorbent assay coated with purified GST-NS1C112aa and immunofluorescence assay with MVCNS1-transfected or MVC-infected WRD cells in chamber slides (Lab-Tek II; Nalge Nunc). Animals with the highest titers were chosen to boost antibody production by injecting 50 μg purified protein through the tail vein. After 2 weeks, boosted animals were euthanized and the terminal blood was collected and the sera harvested. All the animal experiments were approved by the KUMC IACUC.

Immunofluorescence assays were performed essentially as previously described (28).

RESULTS

Sequencing of the palindromic termini of MVC and BPV revealed conserved structures and motifs in the genus Bocavirus. BPV virions are packaged in a way such that 90% of them contain negative-sense DNA and 10% contain positive-sense DNA (12, 13). Therefore, assuming there is a similar maturation mechanism for MVC, we used the strategy of annealing negative- and positive-sense DNA to clone the hairpin structure of MVC. A total of seven clones of the left-end terminus were isolated. Sequencing demonstrated that three were in the flip orientation and four in the flop orientation (Fig. 1A), and they had different end points. Our cloned left-end termini ended at two major sites, nt 1 and nt 18/17, respectively, while one terminus ended at nt 29. The longest left-end terminus had a palindromic structure with 183 nt; similar to BPV, the shortest left-end terminus had only 155 nt (Fig. 1A). The secondary structures of the left-end termini were predicted based on the minimal free energy (ΔG) of the structure. The stem of the MVC left-end terminus contained a “bubble” region and an asymmetric T, which were similar to the features on the left-end hairpin of the minute virus of mice (MVM) (20). In contrast, the BPV left-end stem had three bubble structures, and no unpaired T residues were found (Fig. 1A). The bubble and asymmetric T residue are critical structures in MVM replication of the left-end hairpin (20). Surprisingly, the “rabbit ear” sequences on the left-end terminus of both MVC and BPV were almost identical; only 2 nt at the turn-around region and a pair of G-C dinucleotides were obviously inverted, indicating that sequences of the left-end terminus might share a high degree of conservation in the genus Bocavirus. In other parvoviruses, such as the MVM-like parvoviruses, only the bocaviruses have the distal poly(A) site. It should be emphasized that, among these clones, two had the flop left-end terminus structure and the other two had the flip orientation. Differing ends of both termini were included among the clones. We transfected these four clones into WRD cells, and MVC DNA replication was confirmed by Southern blot analysis with extracted Hirt DNA (Fig. 2A). Newly synthesized DpnI-resistant MVC DNA was detected in cells transfected with all four clones (Fig. 2A, lanes 8, 10, 12, and 14). Interestingly, the monomer ssDNA of MVC either in transfected or in MVC infection was not clearly detectable by Southern blot analysis (Fig. 2A). Similar difficulties in demonstration of ssDNA of AAV5 have previously been encountered; however, ssDNA of BPV was apparently detectable in BPV-infected EBTr cells (J. Qiu, unpublished data). Next, we used cell lysates prepared from the transfected cells to infect WRD cells. At 3 days postinfection, significant cytopathic effects were observed in all four infections (data not shown). MVC infection was confirmed by production of MVC mRNAs (R1 to R5) using the RNase protection assay (Fig. 2B) and by expression of MVC NS1 using the immunofluorescence assay (Fig. 2C). These results demonstrated that all four clones were infectious in WRD cells. Although the presence of different ends of either the left-end or the right-end terminus or different orientations of the left-end terminus (flip or flop) did not make significant differences in the infections, we chose the longest clone (nt 1 to 5402) with the left-end terminus in flip orientation as the infectious clone, named pIMVC throughout this study.
The transcription profiles generated by both MVC infection and transfection of the infectious clone. Total RNA was isolated from MVC-infected WRD cells, and reverse transcription-PCR (RT-PCR), 3' rapid amplification of cDNA ends, and 5' rapid amplification of cDNA ends were performed using various primers. Transcription units shown in Fig. 3B were identified by sequencing the amplified PCR products and confirmed by RNase protection assay (data not shown).

Initially, to determine the relative abundance of various species of MVC mRNA, we assayed both mRNA isolated from MVC-infected cells and pIMVC-transfected WRD cells by Northern blot analysis using the three probes shown in Fig. 3B. The NS probe detected two species of mRNAs at 2.9 kb (R1) and 4.9 kb (R2) (shown in Fig. 3A, lanes 4 and 5, respectively), which were consistent with being unspliced mRNA at the first donor site (1D). Since the first exon was short, mRNAs spliced at 1D were not detected by the NS probe (54). The Cap probe hybridized to mRNAs that ended only at the (pA)d and detected four mRNA bands (Fig. 3A, lanes 2 and 3). They were 4.9-kb (R2), 3.0-kb (R4), 2.5-kb (R5), and 2.2-kb (R6) mRNA. The NSCap probe detected all six major MVC mRNAs, including the 1.0-kb (R3) mRNA that was polyadenylated at the internal polyadenylation site [(pA)p] (Fig. 3A, lanes 6 and 7). The 3.0-kb band of R4 mRNA was barely seen in virus infection (Fig. 3A, lanes 3 and 7), which could be generated less than that in transfection (Fig. 3A, lanes 2 and 6). The overall transcription map suggested by Northern analysis was confirmed by the RNase protection assay (data not shown) and is presented in Fig. 3B. No significant difference was seen between MVC mRNAs generated following infection or transfection of the infectious clone (Fig. 3A, compare lanes 2, 4, and 6 with lanes 3, 5, and 7, respectively). Similar to the map of BPV (54), mid-ORF-encoded mRNAs R3 and R4 were generated that were either polyadenylated at (pA)p or at (pA)d. This is a unique feature of viruses now classified in the genus Bovavirus. A band of mRNA at 3.7 kb that potentially encodes NS2 of BPV was not detected in RNAs isolated from MVC-infected and pIMVC-transfected WRD cells by RT-PCR (data not shown) and Northern blot analysis (Fig. 3A).

MVC NP1 was essential for MVC DNA replication in WRD cells. To test the function of NP1 in MVC infection, we prematurely terminated the mid-ORF by introducing a TAG stop codon at nt 2735 with a single T mutation in pIMVC [NP1 (54)] (Fig. 4B). Transfection of this NP1 knockout mutant, NP1 (54), did not produce significant DpnI-resistant MVC replicative form (RF) DNA (Fig. 4A, lane 5), although NS1 was expressed at a level similar to that from the VP1(54) mutant as seen by immunofluorescence (Fig. 4C). Similar results were also seen when the AUG of NP1 was mutated to ACG, which still left the NS1 ORF unchanged (data not shown). For a comparison, the NS1 ORF was also terminated early in pIMVC [NS1 (54)] (Fig. 4B). Replication of MVC DNA of the NS1(−) mutant...
was totally abolished as shown by immunofluorescence (Fig. 4C). No DpnI-resistant RF DNA was detected (as shown in Fig. 4A, lane 2). Parvovirus NS1 is a multifunctional polypeptide that is essential for parvoviral replication (20). Consistent with this requirement of NS1, our current findings show that MVC NS1 was absolutely required for MVC DNA replication. Surprisingly, NP1 was also essential for MVC DNA replication. Without NP1, replication of MVC DNA was reduced by 320-fold (Fig. 4A, compare lane 1 with lane 5).

To quantify reduction of replication in the absence of NP1, we constructed mutants that could no longer express VP1 or VP2 by mutating the VP1 or VP2 AUG in pIMVC [VP1(H11002)] and VP2(H11002)] (Fig. 4B). DpnI-resistant RF DNA generated from transfection of these two mutants was reduced by approximately five- to eightfold compared with that of the parent infectious clone (Fig. 4A, compare lanes 3 and 4 with lane 1). As VP1 is absolutely required for parvovirus infection (62, 67) and VP2 is the major capsid protein for assembling infectious virions, the VP1 and VP2 knockout mutants did not produce infectious progeny virions (data not shown). Therefore, RF DNA generated from transfection of VP1(−) and VP2(−) mutants resulted only from single-burst replication of transfected plasmid DNA, accounting for the decrease of five- to eightfold in the amount of RF DNA. Compared with the level of RF DNA in transfection of VP1(−) and VP2(−) mutants, transfection of NP1(−) reduced RF DNA by at least 38- to 65-fold (Fig. 4A, compare lane 5 with lanes 3 and 4). This is the second time that a second nonstructural protein of parvoviruses has been demonstrated to be essential for parvoviral monomer form DNA replication (16, 45).

To confirm the function of Bocavirus NP1 in DNA replication, we carried out NP1 cross-complementation tests by providing the NP1 of MVC, BPV, and HBoV in trans, respectively. MVC NP1 increased DNA replication by 8.4-fold, but not to the full wild-type level (Fig. 4A, lane 6). Surprisingly, BPV NP1 and HBoV NP1 were also able to recover MVC RF DNA by 8.8-fold and 5.1-fold, respectively (Fig. 4A, lanes 7 and 8). Expression of these tagged NP1 proteins was confirmed by Western blot analysis in cotransfections (data not shown). These results suggested that functions of Bocavirus NP1 could be cross-complemented among these viruses.

VP1u of MVC contained a phospholipase A2 (PLA2) motif that was critical to MVC infection. To examine the role of the PLA2 motif in viral infection, we mutated three amino acids in the VP1 unique (VP1u) region that are conserved in bocaviruses and in the HDXXY motif of secretory PLA2s (sPLA2s; VP1u35N, VP1u41A, and VP1u42N) (21, 58) (Fig. 5A). For a comparison, two random mutations in the VP1u region, which were outside of the functional PLA2 motif (67), and the nuclear localization sites that are composed of basic amino acid K/R repeats (40, 62, 64) were also made in the infectious clone (VP1u96G and VP1u115P) (Fig. 5A).
These two control mutations were at amino acids 96 and 115. As expected, levels of RF DNA generated in transfection of the PLA2 mutants, VP1u35N, VP1u41A, and VP1u42N, were reduced by approximately eightfold compared with the levels of RF DNA produced by the infectious clone (Fig. 5B, compare lanes 2 to 4 with lane 1). Similarly, the PLA2 mutants achieved the same reduced level of RF DNA as that produced by the VP1(H11002) mutant (Fig. 5B, compare lanes 2 to 4 with lane 7). These results indicated that similar to the RF DNA of the VP1(H11002) mutant, RF DNA of the PLA2 mutants was generated only by single-burst replication of transfected MVC DNA. However, the levels of RF DNA from transfection of the two random mutants, VP1u115P and VP1u96G, of the VP1u region were not significantly reduced (Fig. 5B, compare lanes 2 to 4 with lane 7). These results indicated that the PLA2 motif in VP1u was critical in MVC infection.

DISCUSSION

In this study, we have sequenced the palindromic termini of Bocavirus MVC and have constructed four infectious clones with cloned termini. Systematic analysis of the selected infectious clone revealed common features among viruses in the genus Bocavirus and features unique to parvoviruses. Study of the animal bocaviruses, especially MVC, could help us to elucidate the pathogenesis of HBoV as both MVC and HBoV cause similar symptoms of diseases in their respective hosts (2, 10, 31, 52, 56).

The MVC full-length genome consists of 5,402 nt, and it has disparate palindromic hairpins of 183 nt and 198 nt at the left-end and right-end termini of the genome, respectively. An uncorrected insertion of T at nt 300 was observed in the uncorrected insertion of T at nt 300 was observed in the
previous NSCap sequence (accession no. NC_004442), which extended 58 amino acids at the N terminus of MVC NS1 in the full-length MVC genome (accession no. FJ214110). The fidelity of this MVC genome was confirmed by its infectivity in MVC-permissive WRD cells. Although BPV has been cloned and sequenced, the infectious clone was not available (54). Thus, the infectious clone of MVC reported here will expand our capability to deeply understand the molecular pathogenesis of bocaviruses in general. Moreover, conserved sequences identified in the rabbit ear structures in the left-end terminus and on the stem in the right-end terminus provide information that can be used to clone the termini of HBoV by designing degenerative primers.

The secondary structure of the parvoviral terminus is remarkably important for DNA replication, sometimes more than the sequence per se. It has been shown that the secondary structure of the arm of the AAV terminus is more important than its primary sequences for AAV DNA replication (37). Interestingly, the primary sequence of the rabbit ear structures on the left-end terminus of the two bocaviruses is conserved. The bubble region in the stem of the left-end hairpin BPV was confirmed as being important for BPV DNA replication. Mutation of to-be-paired nucleotides of either the top strand or bottom strand in the bubble region abolished replication in transfected cells (63). The right-end palindromic hairpin of BPV has the capability to form a secondary structure of the stem-arm (Fig. 1B). In another viral model, a similar stem-arm structure that contains the NS1 binding sites had important implications for the viral DNA replication of MVM (6, 18, 19). In comparison, MVC also has the potential to form such a structure of stem-arm at the right-end hairpin (Fig. 1B); however, the free energy in the potential stem-arm structure of MVC is probably not sufficient thermodynamically. The importance of this structure in viral DNA replication warrants further investigation. Interestingly, this stem-arm structure could be created just after the poly(A) signal. It is possible that this structure can slow down RNA polymerase II as a pausing element to terminate transcription for sufficient polyadenylation at the distal poly(A) site (23, 51). Overall, these two bocaviruses, MVC and BPV, shared extensive similarities in structure and conserved motifs on both termini. Identification of these conserved motif sequences may, in the future, be applied to clone the palindromic hairpins of HBoV.

The transcription map of MVC further confirmed that, similar to parvovirus B19 (49) and Aleutian mink disease virus (53), mRNAs from all species of bocavirus were transcribed...
from a single unique promoter and were processed by alternative splicing and alternative polyadenylation (54). We made silent mutations in two actively utilized poly(A) signals in viral infection as well as one AAUAAA signal that was not used in viral infection in the infectious clone. Analysis of RNAs generated by transfection of this mutant showed that MCV RNAs were still polyadenylated at the (pA)p site at a ratio similar to that of the wild type (Y. Sun and J. Qiu, unpublished data). However, these mutations significantly decreased the stabilities of MCV mRNAs, indicating that this region was important to maintain MCV mRNAs in a stable form. Further attempts to knock out the (pA)p sites were not tried since this region is in the coding region for the VP1 unique part.

In all parvoviruses except Aleutian mink disease virus, the minor structural protein, VP1, possesses motifs that have PLAA activity shown to be essential for nuclear entry of the virus during infection (22, 27, 42, 54, 67). The HDXXY motif that was found in the catalytic site of sPLA₂s (21) is also predicted to be present in the VP1u of MVC (58). The aspartic amino acid residue (35D) that is 5 amino acids before the HDXXY motif and the HD amino acids in the HDXXY motif are the most conserved ones in the parvovirus PLAA motifs and sPLA₂s (67). Mutations of these three amino acids in the PLAA motif of the VP1u in the infectious clone confirmed the identity of this protein as VP1, and the PLAA phospholipase motif is present in VP1 that is critical in MVC infection and may play an essential role for virus to access the endosomal membrane (26). Protein analysis of purified MVC by SDS-PAGE showed three structural proteins, VP1, VP2, and VP3, at 81 kDa, 67/63 kDa, and 61 kDa, respectively (58), which was not in agreement with our transcription map of MVC that in only R5 and R6 had capabilities to encode VP1 and VP2. However, three capsid proteins of BPV, VP1 (80 kDa), VP2 (72 kDa), and VP3 (62 kDa), were also detected on SDS-PAGE gels (32, 37). It was suggested that multiple capsid bands might be degraded or cleaved capsid proteins (32, 35). Thus, the nature of these Bocavirus capsid proteins warrants further investigation.

We have demonstrated that MVC NS1, similar to the NS1 of MVM (14), is a multifunctional protein that is essential for viral DNA replication. Parvoviruses express more than one nonstructural protein. The mid-ORF NP1 protein is a bocavirus-unique protein among parvoviruses, which is translated from a continuous ORF located in the middle of the genome. The N terminus of the HBoV NP1 shares the same coding sequence (nt 2537 to 2724) with the C terminus of the NS1 but in a different reading frame. The rest of the coding region of MVC NP1, 369 nt, contained unique coding sequences that lay mostly in the third intron, which was spliced from donor site 3D to acceptor site 3A, in front of the VP1 coding region (Fig. 3B). NP1 has a predicted size of approximately 20 kDa. Similar proteins are not present in other parvoviruses of other genera, setting bacoviruses apart from all other known parvoviruses. Furthermore, NP1 homologous proteins were also not found by BLAST databases. MVC NP1 protein shares an identity of 39.9% and 41.4% with HBoV NP1 and BPV NP1, respectively.

For MVM, the N-terminal exon of NS2 shares its start site and the same reading as NS2 and then after splicing shifts to an alternative ORF (45). The MVM NS2 has a molecular mass of 25 kDa, and the 84-amino-acid N terminus is shared with NS1 and the C terminus is expressed from spliced mRNA with an alternative frame of the NS1-encoding mRNA (17). MVM NS2 is dispensable for productive replication in a variety of nonmurine cells, such as transformed human 324 K cells, but it is required for maximum viral DNA replication in murine cells. The NS2 early-terminated mutants reduced MVM DNA replication maximally by fivefold compared with the wild-type level in a single-burst cycle of replication (16, 45). However, MVC DNA replication was extremely dependent on the function of NP1. An NP1 knockout mutant reduced RF DNA by nearly 50-fold in a single-burst cycle of replication. The NS2 of canine parvovirus (CPV), similar to the MVM NS2, is translated from spliced NS1 ORF and contains 87 amino-terminal amino acids in common with NS1 joined to 78 amino acids from an alternative ORF (65). NS2 knockout mutants in the infectious clone of CPV show no obvious differences in viral DNA replication in cell culture or in dogs (65). However, recent studies examining the within-host genetic diversity of CPV infection showed that NS2 might play a role in host adaptation of CPV (30). MVM NS2 is a multifunctional protein; it also has a role in ssDNA production, capsid assembly, and nuclear export of virions (16, 24, 43, 45, 55). Whether MVC NP1 has multiple roles in MVC infection warrants further investigation. Importantly, BPV NP1 can rescue MVC DNA replication to a similar level as the MVC NP1. Whether the Bocavirus NP1 unique coding sequence can be exchangeable is currently under investigation by replacing the MVC NP1 with BPV or HBoV NP1 in the MVC infectious clone. Thus, we can study the function of HBoV NP1 within the context of MVC.

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REFERENCES

37. Lefebvre, R. B., S. Riva, and K. I. Berns. 1984. Conformation takes prece-


