New, Closely Related Haloarchaeal Viral Elements with Different Nucleic Acid Types

Elina Roine,1⁎† Petra Kukkaro,1†§ Lars Paulin,2 Simonas Laurinavičius,3‡ Aušra Domanska,1|| Pentti Somerharju,3 and Dennis H. Bamford1

Department of Biosciences and Institute of Biotechnology, University of Helsinki, FIN-00014 Helsinki, Finland1; Institute of Biotechnology, DNA Sequencing and Genomics Laboratory, University of Helsinki, FIN-00014 Helsinki, Finland2; and Institute of Biomedicine, Department of Biochemistry and Developmental Biology, University of Helsinki, FIN-00014 Helsinki, Finland3

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During the search for haloarchaeal viruses, we isolated and characterized a new pleomorphic lipid-containing virus, Haloarcula hispanica pleomorphic virus 1 (HHPV-1), that infects the halophilic archaeon Haloarcula hispanica. The virus contains a circular double-stranded DNA genome of 8,082 bp in size. The organization of the genome shows remarkable synteny and amino acid sequence similarity to the genome and predicted proteins of the halovirus HRPV-1, a pleomorphic single-stranded DNA virus that infects a halophilic archaeon Haloarcula sp. Analysis of the two halovirus sequences, as well as the entire nucleotide sequence of the 10.8-kb pHK2-plasmid and a 12.6-kb chromosomal region in Halofexx volcanii, allows us to suggest a new group of closely related viruses with genomes of either single-stranded or double-stranded DNA. Currently, closely related viruses are considered to have the same genome type. Our observation clearly contradicts this categorization and indicates that we should reconsider the way we classify viruses. Our results also provide a new example of related viruses where the viral structural proteins have not diverged as much as the proteins associated with genome replication. This result further strengthens the proposal for higher-order classification to be based on virion architecture rather than on genome type or replication mechanism.

Metagenomic studies have increased the amount of information on the nucleotide sequence space in our environment. It has also increased our awareness of the viral abundance and diversity not recognized before (16, 24, 26). Along with this new information, we have learned to acknowledge the significance of viruses in the evolution and behavior of other organisms (55). To reveal the dynamics and molecular interactions in the interplay between a particular virus and its host, however, isolation of single viruses and their hosts is needed. Even though a number of viruses pathogenic to humans, domestic animals, and plants, as well as some bacteriophages, have been studied in great detail, much of the diversity of the archaeal viruses has remained unknown. By the year 2007 only 44 archaean viruses had been described (2). That embraces less than 1% of all reported viruses. Although the diversity among these few isolated archeal viruses is considerable, a head-and-tail morphology is prevalent among isolated viruses infecting euryarchaeal cells. In contrast, viruses of Crenarchaeota are diverse and often unusual with no viruses having a head-tail morphology (53).

Archeal haloviruses infect euryarchaeal hosts living in environments up to saturated salt. This makes them an interesting group of viruses that reside in a very restricted habitat. In samples taken from high salt environments, the Dead Sea and Spanish solar salterns, viral morphotypes most often observed were spindleshaped, head-and-tail or tailless icosahedral particles (25, 31, 47). Isolated haloviruses, however, do not seem to reflect the proportions of different morphotypes found in the nature as nearly all of the isolates possess a head-and-tail morphology (2). Molecular level studies on only two spindleshaped (10, 11) and one tailless icosahedral particle have been carried out (37, 51). Virus-like particles of other morphologies have also been observed in high-salt environments (47), but only one additional morphotype has been described in detail (50). This recently isolated lipid containing halovirus, HRPV-1, is the first archeal virus containing a single-stranded DNA (ssDNA) genome (50). It infects Halorubrum sp. and has a pleomorphic appearance with glycosylated spike structures protruding from its external membrane (49, 50).

The evolution of prokaryotic viral genome sequences is very fast (18), and the assessment of viral relationships using homology of the genome sequences applies only to closely related viruses (17, 19). Current higher-order classification of viruses is based on the host organism, the nature of the genome (RNA/DNA, single stranded versus double stranded) and the virion morphology. Recently, a higher-order clustering of virus families has been proposed based on common principles of virion architectures as well as on the fold of the major capsid protein...
Viral Elements in Haloarchaea

To determine when virus production begins, a culture was prepared as described above except that an MOI of 1 was used. Samples were drawn at half-hour intervals from 1 h postinfection (hpi) onward for determination of the number of free viruses by plaque assay.

Electron microscopy. For negative stain transmission electron microscopy, viruses were allowed to adsorb to grids for 1 min and then were stained with 1% (wt/vol) potassium phosphotungstate (pH 6.5) for 10 s, uranyl acetate (pH 4.5) for 20 s, 3.32% ammonium molybdate (pH 7.4) for 15 s, or 1% ammonium molybdate (pH 7.0) for 15 s.

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Protein analyses. The protein composition was analyzed by using a modified Tricine-SDS-PAGE system (deviod of spacer gel [59]) with 14% acrylamide concentration (wt/vol). The N-terminal sequences of the two structural proteins were determined in the Protein Chemistry Core Laboratory, Institute of Bio-technology, University of Helsinki, as described previously (8).

Extraction and analysis of the lipids. Lipids from cell suspensions and 2× purified virus preparations were extracted according to the method of Folch et al. (29) essentially as described previously (8) and stored in chloroform-methanol (9:1 [vol/vol]) at −20°C until analyzed. Total lipid extracts were subjected to thin-layer chromatography on silica gel plates (60 F254, Merck) and phospholipids were separated using chloroform–methanol–90% acetic acid (65:35 [vol/vol]) as the solvent (8, 23). After visualizing the lipids by iodine vapor, the plates were sprayed with molybdate (27) and 0.1% orcinol-sulfuric acid reagents (60) to detect phospholipids and glycolipids, respectively. The phosphorus content of lipid spots was determined as described by Bartlett and Lewis (9). The lipids in each spot of the TLC were identified by performing a mass-spectrometric analysis and comparing their m/z values, as well as product and precursor ion profiles, to those of known lipids from halophilic archaea (8).

Genome sequencing and sequence analyses. The genome was purified from the 2×-purified viral material by dilution of the virus 1:50 mMs Tris-HCl (pH 7.2). SDS was added to a final concentration of 1.5%, EDTA was added to 1 mM, and proteinase K was added to 0.8 mg ml−1, and the preparation was incubated at 56°C for 20 min. Proteins were removed by phenol extraction and ether. Nucleic acids were precipitated with 0.3 M sodium chloride and 2 volumes of ethanol at −20°C overnight. Approximately 1.5 to 2 μg of the genome was used for each digest of DNase I (1 U of RQ1 RNase-Free DNase; Promega), RNase A (4 μg; Promega), or mung bean nuclease (0.5 U). As controls, the single-stranded and double-stranded (RFII [RF for replicative form]) viral DNA was purified by cesium chloride (CsCl) 550 ml was measured in hourly intervals of the infected and similarly treated uninfected culture. Samples were drawn concurrently with the absorbance measurements, and the number of free viruses and infective centers was determined by plaque assay. The number of viable cells (CFU/ml) was monitored in both infected and uninfected cultures. After incubation for 12 days, different-sized colonies (small, medium, and large) were picked from viable count plates of the infected culture and inoculated into MGM broth. After incubation at 37°C for 8 days with aeration the titer of the culture supernatants was determined by plaque assay.

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The primers used for PCR and sequencing are listed in Table S6 in the supplemental material. All sequencing was carried out by using BigDye version 3.1 and analyzed on an ABI 3130 (Applied Biosystems). Sequencing reactions were purified with CleanSeq (Agencourt) using a Beckman NXp robot (Beckman Coulter). Sequences were assembled and edited using GAP4 (61). The pHK2 sequence was determined by primer walking using pMDS1 (33) as the template. The primers used for sequence determination are listed in the Table S7 in the supplemental material. The preliminarily annotated sequence of Haloferax volcanii (Hf. volcanii) genome was obtained from the HaloLex website (48).

Genome sequence was analyzed for the open reading frames (ORFs) by using the programs Artemis (56), GenMark (14), and pDRAW32 (AcaClone software). The basic properties of the genes and encoded proteins were studied by using the programs of the EMBOSS program package (54) at the IT Center for Science, Ltd. (www.csc.fi), in Finland and the Expasy proteomics server (30). Transmembrane helices in the polypeptides were predicted by using TMHMM server version 2.0 (41) and coiled coils were predicted using the Coils-Server (45). The global pairwise identity between two proteins was determined by using LALIGN (a global alignment method without end-gap penalty [35] at ch.EMBnet.org) using the default values and the BLOSUM50 matrix. Homologous proteins in the databases were searched by using the BLASTP search engine (3) at the National Center for Biotechnology Information (NCBI).

Nucleotide sequence accession numbers. The nucleotide sequences determined in the present study were deposited in GenBank with the following accession numbers: HHPV-1 genome, GU321093; and pHK2 sequence, GU321094.

RESULTS

**HHPV-1 is a pleomorphic archaeal halovirus.** HHPV-1 was isolated from a solar saltern in Margherita di Savoia, Italy. HHPV-1 produced plaques on the lawn of a *Halorhodopsin* host but was unable to produce plaques on any other strains tested (see Table S1 in the supplemental material).

Halovirus HHPV-1 production and purification methods were established (see Materials and Methods). The virion has densities of 1.27 and 1.23 g/ml in CsCl and sucrose, respectively. Recovery and specific infectivity (PFU/A260) were monitored during the purification process (see Table S2 in the supplemental material). Virions were extremely sensitive to chloroform (data not shown). Chloroform sensitivity and low buoyant density indicated the presence of lipids as virion constituents. Gradient analysis (Fig. 1) confirmed this as *A*, virus titer, proteins, and lipids peaked in the same fractions. Negative-stain transmission electron microscopy (TEM) of HHPV-1 particles purified by equilibrium centrifugation (2× purified virus) showed pleomorphic particles (Fig. 1D).

**HHPV-1 is released without cell lysis.** HHPV-1 did not lyse the host *Halorhodopsin* cells (Fig. 2A). Increase in turbidity was similar for the infected and uninfected cultures until 7 hpi, after which the infected culture showed a slightly retarded growth rate. When the number of CFU was monitored, the infected culture resulted in reduced colony formation after 7 hpi compared to the uninfected one (Fig. 2B).

Virus release started 3 to 3.5 hpi, reaching its maximum, ~8.6×10^{11} PFU/ml in the medium, around 25 hpi (Fig. 2A). The number of infective centers (ICs) started to increase after 3 hpi, indicating that not all of the cells had been infected before the onset of the release of the first viral progeny. Increases of ICs were observed up to ~10 hpi. It should be noted that the amount of ICs could not be detected accurately because viruses seemed to be released continuously from the cells and the detected numbers of ICs were slightly higher than the CFU. Virus production seemed to influence the size of colonies because in the early stages of the life cycle (~1 hpi) colonies formed were mostly large, and cells taken from such colonies did not produce virus. Smaller colonies dominant in the late stages, on the other hand, produced virus.

The life cycle of HHPV-1 was also studied by taking samples for thin-section electron microscopy at different time points during the life cycle. The amount of extracellular viruses was observed to increase in the course of time, and most of the viruses were detected in clusters outside the cells (Fig. 2C). However, we were not able to detect intracellular viruses or viruses in the process of budding.

**There are four phospholipid classes and two major protein species in the HHPV-1 virion.** Analysis of the lipid extracts by TLC revealed that highly purified HHPV-1 virions contain phospholipids but no significant amounts of glycolipids (data not shown). The four phospholipid classes detected were cardiolipins (CL), phosphatidylglycerols (PG), phosphatidylglyc-
erophosphate methyl esters (PGP-Me), and phosphatidylglycerosulfates (PGS). Comparison of the quantity of the three main classes of phospholipids (i.e., PG, PGP-Me, and PGS) of *Har. hispanica*, HHPV-1, and the icosahedral lipid-containing halovirus SH1 (8) shows that the composition of HHPV-1 is quite similar to that of the host, except that the PG is slightly enriched in the virion (Fig. 3), whereas the lipid composition of SH1 deviates significantly from that of the host.

Analysis of the virion in Tricine-SDS-PAGE revealed two major protein species with apparent molecular masses of approximately 60 and 12 kDa (Fig. 4A). The N-terminal sequence analysis of these two proteins gave sequences of AYSNNARGLV for the 12-kDa and VVPVVLPGVA for the 60-kDa protein. These two peptide sequences were identified as the protein products of the ORFs 3 and 4, respectively (see below). The 12-kDa protein is designated VP3 (VP for viral protein), and the 60-kDa protein is designated VP4. The protein sequence analysis also showed that VP4 contains a 50-amino-acid signal sequence that is processed.

**HHPV-1 genome analyses revealed relatedness to the pleomorphic virus HRPV-1.** Enzymatic digestion of the HHPV-1 genome with DNase I and mung bean nuclease showed that the genome is a dsDNA molecule (see Fig. S1 in the supplemental material). The preliminary sequence of the genome was determined by pyrosequencing, and the whole genome sequence was further confirmed by sequencing overlapping PCR-products that covered the whole genome. The ability to generate overlapping PCR-products revealed the circular nature of the genome. The HHPV-1 genome is 8,082 bp in size, and the GC percentage is 55.8%.

The HHPV-1 genome has eight putative ORFs (Fig. 4A). The properties of the ORFs and the encoded polypeptides are shown in Table S3 in the supplemental material. ORFs 3 and 4 encoding the two major structural proteins (Fig. 4A) were designated genes 3 and 4, respectively. The genome organization and protein homology showed that HHPV-1 is closely related to the recently isolated and characterized pleomorphic ssDNA virus HRPV-1 that infects *Halorubrum* sp. (50). The structural protein patterns are also similar (Fig. 4A). HHPV-1 ORF2, and five consecutive translated ORFs show significant identity to the translated ORFs and identified genes of HRPV-1 (Fig. 4). The highest identity (55%) was found between the HHPV-1 gene 3 encoding the small major structural protein VP3 (HHPV-1 VP3) and the VP3 protein of HRPV-1. Although both proteins are putative integral membrane proteins, they differ in the number of predicted transmembrane regions (two in HHPV-1 VP3 and four in HRPV-1 VP3). VP4, the other major structural protein of HHPV-1, is similar in size but shows relatively low identity to the HRPV-1 VP4-protein. Both of these two proteins, however, contain a signal peptide that is processed and a predicted C-terminal transmembrane domain that is preceded by a predicted coiled-coil region.

The remaining putative gene products of HHPV-1 that show homology with the translated ORFs of HRPV-1 are >20% identical (Fig. 4B). The highest identity among these (47%) was found between HHPV-1 ORF7 and the HRPV-1 ORF8. The latter has been shown to encode the minor structural protein VP8 that contains predicted ATPase motifs (50). A search using HHPV-1 ORF7 for conserved domains at the NCBI recognized domains belonging to the P-loop NTPase

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**FIG. 2.** Production of HHPV-1 in a batch culture. (A) The turbidity ($A_{550}$) of infected (●) and uninfected (○) cultures, as well as the number of free viruses (□) were monitored. The time of infection was at 0 hpi. (B) The number of CFU of the infected (■) and uninfected (□) culture. Panels A and B are shown in the same time scale. (C) Thin-section electron micrographs of the infected culture. The number of extracellular viruses grew in the course of time resulting in large clusters of viruses (arrows) at 23 hpi. Bar, 200 nm.

**FIG. 3.** Comparison of the phospholipid compositions of *Har. hispanica*, HHPV-1, and SH1. Concentrations are expressed as the mol% of total phospholipids, and only phospholipids representing >1% of the total are shown. Error bars represent standard deviations of data from at least three independent experiments. PG, phosphatidylglycerol; PGP-Me, phosphatidylglycerophosphate methyl ester; PGS, phosphatidylglycerosulfate.
superfamily proteins (cl09099, Expect [E] value, 0.22). The polypeptide of the predicted ORF1 did not show significant homology to the HRPV-1 ORF1 (9% identity). A BLASTP search of the database at NCBI, however, gives best matches to the putative Rep protein of *Halomicrobium mukohataei* (GenBank accession no. EEJ07478.1, E-value 9e-38) and to ORF1 of plasmid pHK2 (GenBank accession no. AAB09607.1, E-value 2e-25, see below). Since ORF1 of HRPV-1 is also predicted to be a replication initiation protein (50), the two putative ORF1 polypeptides most probably serve the same function as replication initiation proteins.

**Completion of the entire sequence of the pHK2-haloplasmid suggests that it is a virus.** Preliminary BLAST searches using the HHPV-1 ORFs gave hits to the ORFs of the plasmid pHK2 isolated from haloarchaeal species *Haloferax* sp. (34). Since similar observation was made for the HRPV-1 polypeptides (50), we completed the sequence of the pHK2-plasmid using its derivative pMDS1 (33). The previously published sequence of the 4.1-kb fragment of pHK2 (GenBank accession no. L29110.1) (34) was found to diverge from the new sequence in 12 positions. The biggest change was the addition of a G after the nucleotide (nt) 3885 in the previously determined sequence. This change removes the stop codon for ORF4 at nt 3935 and the ORF continues to ORF5. In addition, in the beginning of the previous 4.1-kb region, differences in sequence result in 94% identity between the ORF1 polypeptides predicted to encode replication initiation proteins (data not shown). The sequence obtained from the rest of the pHK2 plasmid showed homologous ORFs in the same order as in both HHPV-1 and HRPV-1 (Fig. 4B; see also Tables S4 and S5 in the supplemental material). The last annotated ORFs, ORF8 of HHPV-1 and ORF9 of HRPV-1, do not show homology to each other or to ORF10 of pHK2 (Fig. 4B). Although the HHPV-1 VP4 shows only 19% identity to the VP4 of HRPV-1, the identity between the pHK2 ORF4 encoded polypeptide and the HRPV-1 VP4 is remarkably higher, i.e., 36%. Also, HRPV-1 VP3 shows higher identity to pHK2 ORF3 (58%) than to HHPV-1 VP3 (Fig. 4B; see Table S6 in the supplemental material). The 10,795-bp pHK2 plasmid has four putative ORFs after the ORF encoding the putative ATPase (HHPV-1 ORF7, HRPV-1 VP8, and pHK2 ORF9). The encoded polypeptides of these ORFs do not show identity to either HHPV-1 or HRPV-1. ORF11, however, encodes a putative polypeptide which shows homology to φH repressor-like proteins and ORF13 encodes a putative integrase (see the supplemental material). These features strongly suggest that...
the pHK2 plasmid actually is a temperate virus related to both HHPV-1 and HRPV-1.

A provirus in the chromosome of *Hfx. volcanii* is the fourth member of the group. A search through preliminarily annotated *Hfx. volcanii* genome using pHK2 ORFs as queries resulted in identification of a 12.6-kb gene region containing ORFs highly identical and in the same order as the ORFs in the pHK2 plasmid (Fig. 4B; see also the supplemental material). The region fulfills the typical features of a provirus integrated into the genome. There are ~450 nt between tRNA^Arg^ and the ORF1 encoding a putative replication initiation protein. ORFs 2 to 9 are 96 to 98% identical to the corresponding ORFs in pHK2; the predicted ATPase encoding ORF9 is the last ORF showing high identity. After that there is no detectable similarity except for ORFs encoding the putative φH repressor-like protein (HVO_1423) and a putative integrase (HVO_1422) that are in similar positions between pHK2 and the chromosomal region of *Hfx. volcanii* in the end of these genetic elements (Fig. 4B). Between the preliminarily annotated ORFs HVO1423 and HVO_1422 there is a short ORF that has not been annotated in the preliminary annotation of the *Hfx. volcanii* genome. The predicted polypeptide, however, shows high homology to the putative ORF12 polypeptide of pHK2.

Apart from this chromosomal region in *Hfx. volcanii* there are several regions encoding hypothetical proteins in the genomes of halophilic archaea that show homology to the viral genomic region starting from VP4 encoding gene and ending after the putative ORF encoding an ATPase. As reported previously (50), the same region also shows homology to the similar region in another halophilic virus, His2 (10).

**DISCUSSION**

During a search for new haloviruses, a pleomorphic halovirus was discovered from a solar saltern on the eastern coast of Italy. This *Har. hispanica* pleomorphic virus 1, HHPV-1, has a pleomorphic appearance similar to the recently reported HRPV-1 virus (Fig. 1D) (50). HHPV-1 production and purification methods were set up to gain highly pure viral material. HHPV-1 virion contains four distinct phospholipid classes and two major structural proteins with predicted transmembrane regions (Fig. 4A). The other described pleomorphic halovirus HRPV-1 has also been shown to contain lipids and two major structural proteins (50). The almost identical phospholipid compositions of HHPV-1 and *Har. hispanica* suggest that lipids during the assembly of HHPV-1 are acquired from the host almost without selection. Interestingly, the composition of phospholipids of SH1, an icosahedral virus that also infects the *Har. hispanica* host, is significantly different from that of HHPV-1 (Fig. 3). This observation indicates that the lipids of SH1 are selectively incorporated into the virion most likely due to the geometrical constraints within the icosahedral protein coat and/or specific lipid-protein interactions (8, 37). Thus, the assembly pathway of HHPV-1 must differ considerably from that of SH1 and related viruses.

HHPV-1 exits the host without causing cell lysis (Fig. 2). Exit without lysis has been observed for several viruses of *Crenarchaeota* (58, 64) and has also been observed for the haloviruses HRPV-1, His1, and His2 (10, 50). Some of these viruses seem to exit in an as-yet-undescribed way. Interestingly, the two viruses homologous to HHPV-1, also exit the host without cell disruption (10, 50). Such a mode of exit suggests budding.

Genome synteny reveals a close relationship between HHPV-1 and the HRPV-1. In addition, the entire 10.8-kb genome of pHK2 completed in the present study as well as the genomic 12.6-kb region of *Hfx. volcanii*, show considerable homology to HHPV-1 and HRPV-1 viruses (Fig. 4B). pHK2 has earlier been reported to be a plasmid (32, 34). Although the viral nature of pHK2 has not been confirmed, the highly homologous genomic region of *Hfx. volcanii* shows the typical features of a virus integrated into tRNA^Arg^.

In addition, both pHK2 and the *Hfx. volcanii* chromosomal region contain a putative integrase (46), as well as genes encoding φH repressor-like proteins. We predict that these two genetic elements can be enclosed into pleomorphic viral particles and consequently will be designated as proviruses in the discussion that follows.

Comparison of the four genomes enables us to detect the most conserved putative and identified proteins of the proposed new group of pleomorphic viruses. Of the two major structural proteins, VP3 and VP4, VP4 shows the highest identity. VP4, on the other hand shows quite high diversity. The VP4 of HRPV-1 was predicted to function as the receptor binding and fusion protein (50). Viral receptor-binding proteins are often the most variable ones among the viral structural proteins (17, 44, 57). Since all of these viruses infect different hosts, variability of VP4 is in accordance with its function as a receptor interacting protein. Apart from these two proteins, ORF5, ORF6, and ORF7 of HHPV-1 show relatively high identity in all four genomes. The region containing identical and almost identical predicted proteins in the pHK2 and *Hfx. volcanii* elements starts from ORF2 and ends after ORF9. In most of the pairwise alignments (see the supplemental material), the predicted and identified proteins of HRPV-1 are more closely related to corresponding ORF-encoded proteins of either pHK2 or *Hfx. volcanii* than to predicted and identified proteins of HHPV-1. HRPV-1 genome also contains ORF5, which shows homology to the corresponding ORFs in pHK2 and *Hfx. volcanii* proviruses. This suggests that HRPV-1 is a closer relative to pHK2 and *Hfx. volcanii* proviruses than to HHPV-1.

Keeping in mind the high identity of the central region of the pHK2 and *Hfx. volcanii* provirus genomes, it is surprising that ORF1 and the region starting from pHK2 ORF10 and *Hfx. volcanii* HVO_1425 varies considerably. HRPV-1 and HHPV-1 genomes do not contain the region starting from the pHK2 ORF10 and including the φH repressor and the integrase. This shows that an integrase is not required for the life cycle of these pleomorphic viruses even though under certain conditions an integrase may confer an advantage. Similar observation have also been made for other types of viruses such as the crenarchaeal virus SSV1 (22) and an icosahedral phage P23-77 that infects *Thermus thermophilus* (38).

The ORFs of the four genomes shows low identity in the pairwise alignments (the highest being 20% between HRPV-1 and *Hfx. volcanii* ORF1). They all are, however, predicted to function as replication initiation proteins. HHPV-1 and pHK2 ORF1 polypeptides contain the conserved domain DUF1424 (E value of < 0.01). This domain can be found in the family of
haloarchaeal proteins predicted to be involved in replication. ORF1-encoded polypeptides of HRPV-1, pHK2, and *Hfx. volcanii* provirus also contain the three conserved motifs suggested to be essential for initiation proteins of rolling-circle replication (36). The HHPV-1 ORF1 contains domains 2 and 3 of these conserved motifs. Since at least HHPV-1, HRPV-1 and pHK2 genomes are circular, it is highly likely that also the ORF1 encodes a rolling-circle replication initiation protein.

Replication and packaging of a circular ssDNA genome of bacteriophage φX174 has been described in detail (40, 63). Replication of the circular ssDNA genome of HRPV-1 and circular dsDNA genome of HHPV-1 may occur in a similar fashion. Packaging of different types of genomes just occurs in different stages of the replication process.

During our studies of new halophilic viruses we have found a new group of pleomorphic viruses including HHPV-1 and HRPV-1 and two putative proviral elements. Based on these results, two major observations can be made that both have consequences as to how viruses are classified. First, we observed that the genome of HHPV-1 is a circular dsDNA molecule, whereas the genome of HRPV-1 is a circular ssDNA molecule (50). In terms of current viral classification, this is unusual, since according to the nucleotide and deduced amino acid sequence data these viruses are closely related and should be grouped together despite the different genome types. This shows that in the era of genome sequencing, higher-order classification according to the genome type may not be justified.

Second, although associated with different hosts, these four viruses show conserved regions, including the two major structural proteins determined for HHPV-1 and HRPV-1 (50). This conserved region containing ORFs 2 to 9 in pHK2 most probably constitutes the core proteins for virion assembly. In contrast, the putative replication initiation proteins show lower identity. In addition, unlike the core proteins, multiple alignment of the ORF1 polypeptides arranges the relationships between the viruses and the proviral elements in a different order than the alignment using the core proteins. Thus, comparison of the four genomes suggests that the viral world is indeed “… a dynamic network of relationships in which genes have diverse, variously intertwined histories …” (39) obtained both vertically and horizontally. The comparisons of the core proteins of this new pleomorphic virus group show, however, that the structural components are more conserved and most probably obtained by vertical inheritance in order to keep the viral self intact and functional for continuation of the viral lineage (1, 4–6, 12, 13, 42). Similar observations have been made for tailed phages using even more abundant sequence information (20, 21, 28, 62). As a consequence, the mode of the genome replication and the nature of the genome (RNA/DNA, ss or ds) may not be adequate determinants for classifying distant, but related viruses to higher-order taxa. The determinants for the virion architecture instead may better serve this function.

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