

# Three-Dimensional Structure of a Protozoal Double-Stranded RNA Virus That Infects the Enteric Pathogen *Giardia lamblia*

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#### **ABSTRACT**

Giardia lamblia virus (GLV) is a small, nonenveloped, nonsegmented double-stranded RNA (dsRNA) virus infecting Giardia lamblia, the most common protozoan pathogen of the human intestine and a major agent of waterborne diarrheal disease worldwide. GLV (genus Giardiavirus) is a member of family Totiviridae, along with several other groups of protozoal or fungal viruses, including Leishmania RNA viruses and Trichomonas vaginalis viruses. Interestingly, GLV is more closely related than other Totiviridae members to a group of recently discovered metazoan viruses that includes penaeid shrimp infectious myone-crosis virus (IMNV). Moreover, GLV is the only known protozoal dsRNA virus that can transmit efficiently by extracellular means, also like IMNV. In this study, we used transmission electron cryomicroscopy and icosahedral image reconstruction to examine the GLV virion at an estimated resolution of 6.0 Å. Its outermost diameter is 485 Å, making it the largest totivirus capsid analyzed to date. Structural comparisons of GLV and other totiviruses highlighted a related "T=2" capsid organization and a conserved helix-rich fold in the capsid subunits. In agreement with its unique capacity as a protozoal dsRNA virus to survive and transmit through extracellular environments, GLV was found to be more thermoresistant than Trichomonas vaginalis virus 1, but no specific protein machinery to mediate cell entry, such as the fiber complexes in IMNV, could be localized. These and other structural and biochemical findings provide a basis for future work to dissect the cell entry mechanism of GLV into a "primitive" (early-branching) eukaryotic host and an important enteric pathogen of humans.

#### **IMPORTANCE**

Numerous pathogenic bacteria, including Corynebacterium diphtheriae, Salmonella enterica, and Vibrio cholerae, are infected with lysogenic bacteriophages that contribute significantly to bacterial virulence. In line with this phenomenon, several pathogenic protozoa, including Giardia lamblia, Leishmania species, and Trichomonas vaginalis are persistently infected with dsRNA viruses, and growing evidence indicates that at least some of these protozoal viruses can likewise enhance the pathogenicity of their hosts. Understanding of these protozoal viruses, however, lags far behind that of many bacteriophages. Here, we investigated the dsRNA virus that infects the widespread enteric parasite Giardia lamblia. Using electron cryomicroscopy and icosahedral image reconstruction, we determined the virion structure of Giardia lamblia virus, obtaining new information relating to its assembly, stability, functions in cell entry and transcription, and similarities and differences with other dsRNA viruses. The results of our study set the stage for further mechanistic work on the roles of these viruses in protozoal virulence.

"iardia lamblia (also called G. intestinalis or duodenalis) is a Iflagellated, parasitic protozoan and an important cause of diarrheal disease worldwide, especially in developing countries. It is a member of the supergroup Excavata, one of the earliest diverging eukaryotic taxa, to which the parasitic protozoa Leishmania, Naegleria, Trichomonas, and Trypanosoma also belong, although in distinct groups or subgroups relative to Giardia (1). Infection by G. lamblia occurs after ingestion of cysts from contaminated water, food, or soil or sometimes by direct contact. After excystation and multiplication, G. lamblia trophozoites adhere to the surface of epithelial cells in the small intestine. Specific pathogenic mechanisms remain poorly characterized, but the resulting diarrheal disease is characterized by malabsorption, maldigestion, chloride hypersecretion, and increased intestinal transit rates (2, 3). Histopathological findings include lymphocyte infiltration and diffuse blunting of the microvilli, enterocyte apoptosis, and disruption of epithelial tight junctions.

Epidemiologically, *G. lamblia* is the most common protozoan pathogen of the human small intestine, estimated to cause ≥280 million cases yearly (2, 4), and is also common in companion animals and livestock. Most infections are self-limiting, but in

immunocompromised persons, including undernourished children and people with AIDS, *G. lamblia* disease may be more chronic, severe, and atypical and may be associated in children with growth retardation and cognitive impairment (5, 6). Nota-

Received 22 September 2014 Accepted 31 October 2014 Accepted manuscript posted online 5 November 2014

Citation Janssen MEW, Takagi Y, Parent KN, Cardone G, Nibert ML, Baker TS. 2015. Three-dimensional structure of a protozoal double-stranded RNA virus that infects the enteric pathogen *Giardia lamblia*. J Virol 89:1182–1194. doi:10.1128/JVI.02745-14.

Editor: S. López

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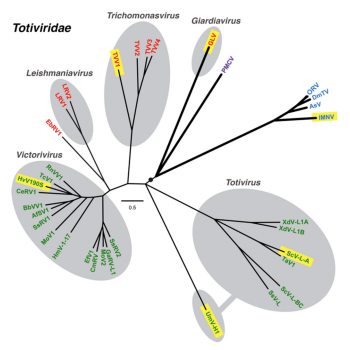


FIG 1 Phylogenetic relationships among taxonomically approved and tentatively assigned members of family Totiviridae. A maximum-likelihood phylogenetic tree was reconstructed from sequences encompassing the predicted CP+RdRp products of each analyzed virus, as described in Materials and Methods. Branches with support values <70% were collapsed to the preceding node, and some terminal branches within the genus Victorivirus are not labeled for simplicity but were all supported at ≥97%. Viruses are labeled with abbreviated names (see Materials Methods for key and GenBank accession numbers), and those for which 3D structures have been determined are highlighted in yellow. Protozoal viruses are labeled in red, fungal viruses are labeled in green, arthropod viruses are labeled in blue, and the one vertebrate virus is labeled in purple. Approved Totiviridae members are grouped within gray ovals according to current genus assignments as labeled; fungal virus UmV-H1 is currently grouped in genus *Totivirus* as shown but is divergent and appears to warrant reclassification. The black circle and thicker branch lines indicate an apparent subclade of viruses that can undergo efficient extracellular transmission. The scale bar indicates the number of substitutions per aligned position.

bly, like several other protozoan pathogens, *G. lamblia* can undergo antigenic variation to evade host adaptive immunity (7), contributing to its capacity for chronic infections. It can also occur in outbreaks due to localized contamination of food or water sources and is therefore recognized by the National Institutes of Health/National Institute of Allergy and Infectious Disease (NIH/ NIAID) as a category B priority pathogen (http://www.niaid.nih.gov/topics/biodefenserelated/biodefense/pages/cata.aspx) associated with food and waterborne illness.

G. lamblia itself is susceptible to infection by a small, nonenveloped double-stranded RNA (dsRNA) virus called Giardia lamblia virus (GLV). Discovered in the 1980s (8), GLV represents the prototype and indeed is the sole species in the genus Giardiavirus, family Totiviridae (Fig. 1). It has a nonsegmented, 6.3-kbp genome with two, long, partially overlapping open reading frames (ORFs) on one of the genomic RNA strands, respectively encoding the capsid/coat protein (CP; ORF1) and the RNA-dependent RNA polymerase (RdRp; ORF2) (9). In addition, the ORF1 product undergoes proteolytic cleavage(s) to remove its 32 N-terminal amino acids, which appear not to be present in mature virions (10). The RdRp is expressed in fusion with the CP (i.e., as

CP/RdRp) by means of -1 ribosomal frameshifting in the region of ORF1-ORF2 overlap (9, 11). The GLV mRNA, which spans the full length of the genomic plus strand, lacks a 5' cap and is not polyadenylated; translation is instead initiated by means of an internal ribosomal entry site that partially overlaps ORF1 (12, 13).

Several other parasitic protozoa, including *Leishmania*, *Trichomonas*, and *Cryptosporidium*, are also known to carry small dsRNA viruses: *Leishmania* RNA viruses (LRVs) 1 and 2 (genus *Leishmaniavirus*, family *Totiviridae*); *Trichomonas vaginalis* viruses (TVVs) 1, 2, 3, and 4 (genus *Trichomonasvirus*, family *Totiviridae*); and *Cryptosporidium parvum* virus 1 (genus *Cryspovirus*, family *Partitiviridae*). Moreover, several types of evidence suggest that these other protozoal dsRNA viruses might enhance the pathogenicity of their respective protozoa for causing mammalian disease (14–19). It is not yet clear whether the same may be true for GLV in exacerbating *Giardia*-associated disease, although an early review by Wang and Wang (20) indicated there is no obvious correlation between GLV and *Giardia* virulence.

Interestingly, GLV is the only one of the known protozoal dsRNA viruses that can transmit efficiently by extracellular means (8, 21), meaning that GLV virions uniquely include the protein machinery for mediating efficient cell entry. The GLV-Giardia virus-host system thus provides an unusual opportunity to determine the steps involved in cell entry of a "primitive" (early-branching) eukaryote, with a simplified endo/lysosomal vacuolar system (22, 23). Attachment of GLV to the *G. lamblia* host cell has been shown to involve a cell surface receptor, which is missing from some *G. lamblia* strains (21, 24), but the identity of this receptor remains unknown. Otherwise, the entry mechanism of GLV is poorly characterized, except for one report suggesting that it is susceptible to inhibition by ammonium chloride and chloroquine and thus may require endocytosis and vacuolar acidification (25).

Resemblance of GLV to other Totiviridae members has not been previously confirmed at the structural level, except by negative-stain transmission electron microscopy (TEM) showing nonenveloped particles with single-layered capsids, appearing to measure 340 to 370 Å in diameter (8, 26). Moreover, in addition to its virions containing the protein machinery for efficient cell entry, which the other Totiviridae members from protozoal or fungal hosts do not, GLV appears to be more closely related to a group of nonsegmented dsRNA viruses isolated in recent years from metazoan hosts, both arthropods (crustaceans and insects) and vertebrates (bony fish) (27-31) and tentatively also assigned to family Totiviridae (Fig. 1). Notably, these metazoan viruses share with GLV the ability to transmit efficiently by extracellular means. The virions of one of these viruses, penaeid shrimp infectious myonecrosis virus (IMNV), have been analyzed by transmission electron cryomicroscopy (cryo-TEM) and icosahedral three-dimensional (3D) image reconstruction, which revealed an unusual proteinfiber complex that projects above the so-called "T=2" capsid at the icosahedral 5-fold axes and is likely to play one or more key roles in cell entry (32, 33). Such "T=2" capsid organization (34) reflects a T=1 structure in which a CP dimer occupies each of the icosahedral asymmetric units (120 CP subunits per capsid) and is common to many dsRNA viruses, including other members of family Totiviridae (35–39).

Considering the variety of interesting biological aspects of GLV and also the importance of its protozoan host *G. lamblia* as a

human disease agent, we undertook this study to determine the virion structure of GLV via cryo-TEM and icosahedral 3D image reconstruction. In parallel, we also investigated several of the *in vitro* biochemical properties of GLV virions. The results of these studies reveal both similarities and differences between GLV and other *Totiviridae* members, including TVV1, prototype fungal totivirus *Saccharomyces cerevisiae* virus L-A (ScV-L-A), and tentatively assigned member IMNV, and set the stage for ongoing studies of cell entry by GLV.

## **MATERIALS AND METHODS**

Protozoan cultures and virion purifications. A frozen aliquot of G. lamblia strain WBI (GLV-infected WB) cells was kindly provided by C. C. Wang and coworkers (University of California–San Francisco). The batch culture was maintained as described previously (26) and expanded in cell numbers and volume over several days before harvest for virion purification. At harvest time, containers were chilled in an ice bath and swirled to remove attached cells from the glass surface. GLV virions were customarily purified from a batch of  $2 \times 10^9$  WBI cells, using the same protocol of sonication and centrifugations as previously described for TVV1 virions (18, 39). GLV virions obtained from the last centrifugation step (banding in a CsCl density gradient) were dialyzed overnight against HNM buffer (50 mM HEPES [pH 7.2], 500 mM NaCl, 20 mM MgCl<sub>2</sub>) and then kept at  $4^{\circ}$ C for up to 3 weeks or at  $-80^{\circ}$ C for longer periods before use. Concentrations of the dialyzed preparations routinely measured near 200 µg/ml (~10<sup>12</sup> virions/ml) according to a modified Bradford protein assay (Bio-Rad). T. vaginalis isolate UH9 was cultured, and TVV1-UH9 virions were purified as previously described (18, 39). The original T. vaginalis UH9 stock, which was kindly provided by Raina N. Fichorova and coworkers (Brigham and Women's Hospital and Harvard Medical School), had been serially cloned twice in soft agar before use in this study.

Liquid chromatography-tandem mass spectrometry (LC-MS/MS). Two aliquots of purified GLV virions (each  $\sim 2 \times 10^{10}$  in 20  $\mu$ l of HNM buffer) were denatured by heating to 80°C for 5 min and submitted to the Beth Israel Deaconess Medical Center Mass Spectrometry Facility for processing and analysis. Separate digestions were performed with sequencing-grade stocks of trypsin or chymotrypsin. Resulting spectra were analyzed using the Sequest search engine (40) against a customized database of GLV CP and CP/RdRp sequences. Analyses were otherwise performed as described previously (39).

TEM. Prior to performing cryo-TEM, the integrity and homogeneity of samples were evaluated by using negative-stain TEM. For this, 3.5- $\mu l$  aliquots of sample were absorbed to a continuous carbon grid that had been glow-discharged for  $\sim\!60\,s$  in an Emitech K350 evaporation unit. The absorbed sample was then stained with 1% (unbuffered) uranyl acetate and rinsed with deionized distilled water. Micrographs were recorded on a 4K² Gatan Ultrascan charge-coupled device camera in an FEI G2 Tecnai (Polara) microscope. Initial experiments performed to judge proper storage conditions for purified GLV virions and to provide an initial gauge of preparation quality were performed similarly, but using uranyl formate as stain and a JEOL 1200EX 80-kV microscope at the Harvard Medical School Department of Cell Biology Conventional EM Facility.

For cryo-TEM, 3.5- $\mu$ l aliquots of sample were vitrified and examined using established procedures (36). In brief, samples were applied to Quantifoil holey grids that had been glow-discharged for  $\sim\!15$  s. Grids were then blotted with Whatman filter paper for  $\sim\!5$  s, plunged into liquid ethane, and transferred into a precooled FEI Polara multispecimen holder, which maintained the specimen at liquid-nitrogen temperature. Micrographs were recorded on Kodak SO-163 electron-image film in the Polara microscope operated at 200 keV and under minimal-dose conditions ( $\sim\!25$  e/A²) at a nominal magnification of  $\times\!59,000$  and with the objective lens settings ranging between 0.78 and 3.41  $\mu$ m underfocus.

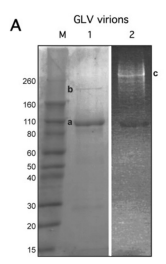
**Icosahedral 3D image reconstruction.** Micrographs exhibiting minimal astigmatism and specimen drift were digitized at 6.35-μm intervals (1.09-Å pixel size at the specimen) on a Nikon Super Coolscan 8000 mi-

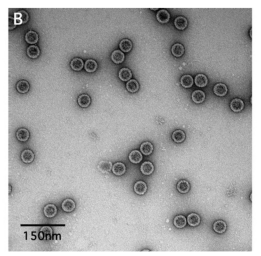
crodensitometer. RobEM (http://cryoem.ucsd.edu/programs.shtm) was used to extract 14,125 particle images, each  $601 \times 601$  pixels in size, and to preprocess the images as described previously (41). The defocus and astigmatism levels of each micrograph were estimated using the program CTFFIND3 (42). The random-model computation procedure (43) was used to generate an initial 3D density map at an  $\sim$ 29-Å resolution from the images of 150 particles. This map served as a starting model to initiate full orientation and origin determinations of the entire set of images using AUTO3DEM v4.03.1 (44). After several cycles of orientation and origin refinement with the PO²R and P3DR functions of AUTO3DEM, the resolution converged to  $\sim$ 6.0 Š(based on the 0.5 threshold of the Fourier shell correlation criterion [45]). The effects of the microscope contrast-transfer function were compensated in part by inverting image phases in spatial frequency bands where contrast reversals occurred (46). An inverse temperature factor of 1/(80 Ų) was applied to the final reconstruction (47).

The Segger v1.7 tool in the Chimera visualization package (48) was used to segment out individual CP-A and CP-B subunits in the final map and to examine interactions between them as described previously (38). Briefly, segmentation is obtained with this tool by means of an iterative procedure, starting from an initial arbitrary assignment of density regions to the two subunits and then refining those assignments in a semiautomatic manner by imposing additional restrictions. The additional restrictions in this case were based on evidence from the estimated capsid volume that the capsid indeed comprises 120 CP subunits, i.e., two CP subunits per icosahedral asymmetric unit, and on the following assumptions: (i) each subunit forms a compact domain, (ii) global morphology and particular features are repeated in CP-A and CP-B, (iii) symmetry constraints of the icosahedral arrangement of subunits within the capsid are maintained, and (iv) no obvious capsid density is left unassigned. Under these constraints, the final result does not depend on the initial arbitrary assignment to the two subunits. GLV subunit A was manually docked into GLV subunit B for comparison and the Chimera "fit-in-map" tool was then used to refine the fit. The normalized correlation value of the final fitting was 0.91 as provided by Chimera (48). Location and length of density features that could be assigned to putative  $\alpha$ -helical segments in the segmented A subunits of the different totiviruses were predicted automatically using the VolTrac algorithm (49) implemented in the Sculptor graphics software (50). The procedure was performed as described by Dunn et al. (38).

Temperature stability of virions. GLV or TVV1 virions ( $2\times10^9$  to  $5\times10^9$  in  $10~\mu$ l) were incubated in HNM buffer for 5 min at the indicated temperatures, using the gradient setting of a thermal cycler (MultiGene; Labnet International). RNase III (New England BioLabs) was then added to a final concentration of  $0.02~\text{U/\mu l}$ , followed by incubation for 5 min at 37°C to digest dsRNA that was released from, or made accessible inside, destabilized virions. RNase III digestion was quenched by addition of EDTA to a final concentration of 100 mM. Protected dsRNA was then released from intact virions by heating the mixture for 5 min at 65–75°C. The genomic dsRNA bands of GLV and TVV1 were visualized on a 0.7% agarose gel, stained with ethidium bromide, and quantitated by using ImageJ software (http://imagej.nih.gov/ij/). The plot and fit for Fig. 6A were generated using DataGraph 3.1.1.

In vitro transcription. GLV or TVV1 virions ( $10^8$  in  $20~\mu$ l) were incubated in transcription buffer (150~mM NaCl; 2~mM MgCl;;  $200~\mu$ M concentrations each of ATP, CTP, and UTP;  $20~\mu$ M [ $\alpha$ - $^{32}$ P]GTP; and 50~mM Tris-HCl or Tris/acetate buffer at the indicated pH value) for 30~min at  $37^\circ$ C. The transcription reaction was terminated by addition of formamide and EDTA to final respective concentrations of 30% and 17~mM. Transcripts were separated on an 8% denaturing polyacrylamide gel containing 7~M urea in  $0.5\times$  Tris-borate-EDTA buffer. Radiolabeled RNA bands were then visualized by phosphorimaging on a Typhoon 9400~variable mode imager and quantitated using ImageQuant TL software (GE Healthcare). The plot and fit for Fig. 6B were generated using DataGraph 3.1.1.





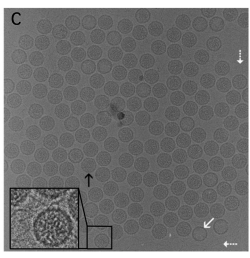


FIG 2 Gels and electron micrographs of purified GLV virions. (A) Gel analyses. SDS-PAGE and Coomassie blue staining (left) shows the GLV capsid protein (a) migrating near 100 kDa and the GLV CP/RdRp (b) near 190 kDa. Minor protein bands near 85 and ~28 kDa are thought to be contaminants or degradation products. Agarose gel electrophoresis and ethidium bromide staining (right) shows a single band for GLV dsRNA (c). (B) Negative-stain TEM. Virions were stained with uranyl acetate to assess particle quality. The scale bar also applies to panel C. (C) Cryo-TEM. Unstained, vitrified virions are shown in one field used for image reconstruction. Examples of full and empty GLV particles are highlighted with black and white arrows, respectively. White dotted arrows point to putative strands of free dsRNA in the solvent background, also seen in the enlarged inset at higher contrast.

Phylogenetic tree and polarity plot. A maximum-likelihood phylogenetic tree was reconstructed from sequences encompassing the predicted CP+RdRp products of each analyzed virus (see Fig. 1). Multiplesequence alignments were conducted using MAFFT 7.1, as implemented with default settings at http://mafft.cbrc.jp/alignment/software. Model testing was then performed using ProtTest 3.3 (51), which showed the Blosum62 substitution matrix and I+G+F distributions (proportion of invariable sites, 0.01; gamma shape parameter, 2.975; and empirically determined amino acid frequencies) to score most highly according to both Akaike and Bayesian information criteria. Phylogenetic analyses were next performed with these predetermined parameters using PhyML 3.0 (52) as implemented at http://www.hiv.lanl.gov/. The starting tree was optimized by both branch length and tree topology, and tree improvement was performed according to the best of nearest neighbor interchange and subtree pruning and regrafting. Branch support values (%) were estimated by the approximate likelihood ratio test with SH-like supports. The final tree was refined for display using FigTree 1.4.0. The abbreviations (along with the corresponding GenBank accession numbers) for the viruses included in the tree are as follows: GLV (NC 003555); TVV1 (NC\_003824); TVV2 (NC\_003873); TVV3 (NC\_004034); TVV4 (HQ607522); LRV1 (NC\_002063); LRV2 (NC\_002064); EbRV1, Eimeria brunetti RNA virus 1 (NC\_002701); SsRV2, Sphaeropsis sapinea RNA virus 2 (NC\_001964); GaRV1-L1, Gremmeniella abietina RNA virus L1 (NC\_003876); MoV2, Magnaporthe oryzae virus 2 (NC\_010246); CmRV, Coniothyrium minitans RNA virus (NC\_007523); EfV1, Epichloe festucae virus 1 (AM261427); HmTV1-17, Helicobasidium mompa totivirus 1-17 (NC\_005074); MoV1, Magnaporthe oryzae virus 1 (NC\_006367); SsRV1, Sphaeropsis sapinea RNA virus 1 (NC\_001963); AfSV1, Aspergillus foetidus slow virus 1 (HE588147); BbVV1, Beauveria bassiana victorivirus 1 (HE572591); RnVV1, Rosellinia necatrix victorivirus 1 (AB698489); CeRV1, Chalara elegans RNA virus 1 (NC 005883); HvV190S, Helminthosporium victoriae virus 190S (NC\_003607); TcV1, Tolypocladium cylindrosporum virus 1 (NC\_014823); XdV-L1A, Xanthophyllomyces dendrorhous virus L1A (NC\_020903); XdV-L1B, Xanthophyllomyces dendrorhous virus L1b (JN997473); TaV1, Tuber aestivum virus 1 (HQ158596); ScV-L-A (L1) (NC\_003745); ScV-L-BC (La) (NC\_001641); SsV-L, Scheffersomyces segobiensis virus L (KC610514); UmV-H1, Ustilago maydis virus H1 (NC\_003823); ORV, Omono River virus (AB555544); DmTV, Drosophila melanogaster totivirus (NC\_013499); AsV, Armigeres

subalbatus virus (NC\_014609); IMNV (NC\_007915); and PMCV, piscine myocarditis virus (NC\_015639).

A polarity plot was generated for the full-length GLV ORF 1 product (CP precursor) according to the method of Grantham (53) as implemented at http://web.expasy.org/protscale/ with a running window size of 15 (see Fig. 3B).

Accession number. The 3D density map of the cryo-reconstruction of the GLV virion has been deposited in the Electron Microscopy Data Bank at the European Bioinformatics Institute with accession code EMD-5948.

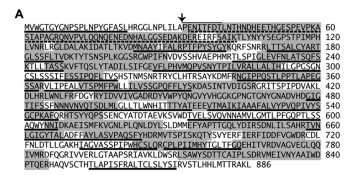
# **RESULTS**

# Initial characterizations of virions used for cryo-TEM studies.

G. lamblia strain WBI (strain WB infected with GLV strain Portland I) was analyzed, because it has been extensively studied by Wang and coworkers (8–14, 21, 24–26). WBI was serially passaged in liquid batch culture, with a doubling time of  $\sim$ 24 h. After harvest, cells were sonicated to release virions, centrifuged to deplete debris, and sedimented into a CsCl density gradient. The gradient was fractionated, and fractions containing virions were identified by SDS-PAGE to track GLV CP. Virions consistently concentrated in only one or two consecutive fractions, corresponding to a visible band in the gradient, well separated from any debris. The virion fractions were dialyzed against buffer and stored at either 4°C or -80°C before use.

SDS-PAGE of the purified GLV virions showed a major Coomassie-stained band migrating near 100 kDa, a finding consistent with the sequence-predicted mass of 95 to 98 kDa for GLV CP (95 kDa if missing its N-terminal 32 amino acids [aa] [10]) (Fig. 2A). A minor Coomassie blue-stained band migrating near 200 kDa was also routinely visible, consistent with the lower copy number and sequence-predicted mass of 211 kDa for GLV CP/RdRp (only 1 or 2 copies per virion versus 119 or 118, respectively, for CP). An ethidium-stained band migrating near the top of the SDS-PAGE gel, and consistent with the GLV dsRNA genome, was additionally seen (Fig. 2A).

To evaluate suitability of the sample for structural analysis, we



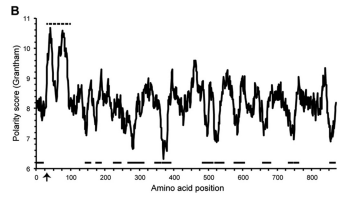


FIG 3 Mass spectrometry to assess protein contents of GLV virions. (A) Sequence of GLV CP precursor is shown. Arrow indicates position of the N-proximal processing cleavage (overlining, N-terminal sequencing results [10]). Sequences represented in the peptides recovered from LC-MS/MS after trypsin or chymotrypsin cleavage of GLV virions are indicated by gray shading. Predominantly uncharged regions as defined in the Discussion are underlined. The highly charged region near the CP N terminus is indicated by dotted underlines. (B) A polarity plot was generated as described in Materials and Methods. Highlighted features (arrow, underlines) are translated from panel A.

negatively stained the purified GLV virions and inspected them by TEM. The particles were seen to be well distributed, with minimal breakage and consistent widths near 450 Å (Fig. 2B). Most displayed a regular, circular outline. Essentially all particles appeared to be penetrated by stain, but most had a heterogeneous distribution of interior staining consistent with the presence of centrally packaged GLV genome. A few "empty" particles, with more homogeneous interior staining consistent with the absence of genome, were also observed. Similar findings were obtained for purified virions that had been stored at 4°C or at -80°C for several weeks, with or without addition of glycerol. Storage at -20°C was less reliable, however, as particles in clumped, empty, and broken states were observed in increased numbers.

Purified GLV virions were submitted for LC-MS/MS to assess their protein contents. Parallel analyses were performed: one for peptides generated by trypsin treatment and the other by chymotrypsin treatment. The identified peptides, respectively, covered 56 and 46% of the deduced CP sequence, or 65% coverage by the combined results (Fig. 3A). A notable finding was that no peptides were identified from the N-terminal 32 aa of the ORF1-encoded CP sequence, which have been indicated by Yu et al. (10) to be removed by proteolytic cleavage in infected cells and to be missing from virions. Peptides were also recovered from the RdRp region of CP/RdRp in these analyses, although at lower frequency, 25%

coverage of the predicted RdRp sequence by the combination of peptides from either treatment, consistent with the lower copy number of CP/RdRp. Peptides spanning the CP/RdRp fusion junction were not recovered.

Cryo-TEM and 3D reconstruction: surface views and equatorial section. Purified GLV virions vitrified and visualized by cryo-TEM appeared consistently sized and shaped, with mostly unbroken capsids (Fig. 2C). The genomic dsRNA enclosed within the capsid layer appeared as distributed density, without the "fingerprint" motif that is observed in virions of other totiviruses, including tentatively assigned member IMNV (32, 35, 36, 38, 39). A few particles, however, exhibited lower central densities, and linear strands suggestive of free dsRNA were seen at places in the solvent background, consistent with loss of genome from a few particles in the preparation (Fig. 2C). The capsids in this mixture of mostly "full" and few "empty" particles displayed a very regular, circular outline, without any obvious protrusions. An icosahedral 3D image reconstruction was computed from 14,125 particle images recorded on 101 micrographs. The resolution of the final reconstruction was estimated at 6.0 Å based on a Fourier-shell correlation threshold of 0.5 (45).

As expected for totiviruses (32, 35–39), a surface view of the 3D reconstruction revealed that GLV appears to possess a so-called "T=2" capsid (34) comprising 120 CP subunits in a T=1 lattice (Fig. 4A and C). The estimated volume of the capsid shell is also consistent with this conclusion. Large portions of the CP outer surfaces form a raised but mostly flat "plateau" centered on each of the icosahedral 5-fold axes, with the portion of each subunit contributing to this plateau having the appearance of an elongated, triangular "pizza slice." The wider ("crust") end of each slice lies distal to the 5-fold axis. From this perspective, it is notable that each CP subunit occupies either of two nonidentical positions within the capsid (Fig. 4A). In one position (designated "A" as in other "T=2" capsids), the CP subunits approach one another around each 5-fold axis via the narrow ("toppings") end of the figurative slice, while in the other position (designated "B"), the CP subunits lie wedged between neighboring A subunits, in a similar orientation but more distant from the 5-fold axis. Each plateau thus appears to be constituted by portions of 10 CP subunits: five A and five B. By comparison, each of the icosahedral 3-fold axes, which are low points on the capsid surface, appears to be immediately surrounded by three B subunits, each from a different decameric plateau (Fig. 4A). Each of the icosahedral 2-fold axes, on the other hand, appears to be immediately surrounded by four CP subunits: one A and one B subunit from one decameric plateau and one A and one B subunit from the adjacent decameric plateau (Fig. 4A). Moreover, each of these A subunits appears to make contact in a quasi-2-fold-symmetrical, "tail-to-tail" (or "crust-to-crust") fashion with the B subunit from the adjacent decamer, thereby forming two elongated A:B ridges that pass alongside the icosahedral 2-fold axis, which is another low point on the capsid surface.

The equatorial section of the 3D reconstruction shows both punctate and linear features, which at this level of resolution suggest a prevalence of  $\alpha$ -helical secondary structure in the CP subunits (Fig. 4B). It also shows that the single-layered capsid of GLV has a maximum thickness of  $\sim 60$  Å (median,  $\sim 40$  Å). The outermost capsid diameter is  $\sim 485$  Å, making this the largest totivirus capsid studied to date (35–39), including the capsid (sans spikes) of IMNV (32). The central section also reveals an interior capsid

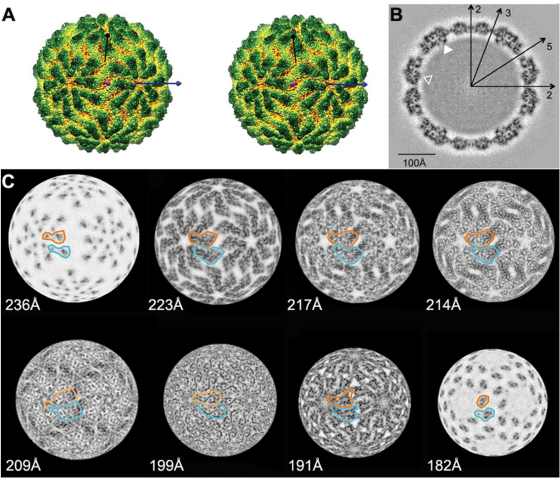


FIG 4 Icosahedral 3D image reconstruction of GLV virions. (A) Space-filling stereo view of the GLV particle surface as viewed down an icosahedral 2-fold axis (magenta). One icosahedral 3-fold axis (black) and one icosahedral 5-fold axis (blue) are marked. The map is color-coded by radius (dark green, outermost; red, innermost). (B) Central (equatorial), 1-pixel (1.09-Å)-thick section through the virion density map shown in grayscale (black and white corresponding to highest and lowest densities, respectively). Symmetry axes (icosahedral two-, three-, and 5-fold) are marked, as well as close contacts between capsid and RNA densities (arrowheads). (C) Radial sections through the capsid, centered at indicated radii and each 1 pixel (1.09 Å) thick. The map is coded in grayscale as in panel B. The outlines of one CP-A (orange) and one CP-B (cyan) subunit are shown in each radial section.

surface that is smoother than the exterior, except for short inward protrusions near the icosahedral 3-fold axes.

Unlike what has been observed to date for other totiviruses (32, 35, 36, 38, 39), the packaged genome of GLV does not appear in the reconstruction as a series of concentric shells of density in the particle interior. This finding is consistent with the lack of central fingerprint densities in the raw cryo-micrographs of GLV (Fig. 2C). Instead, densities attributable to genome are distributed almost homogeneously throughout the interior of the capsid, except for a hint of greater density in some localized regions nearest the shell (Fig. 4B). The most notable contacts between capsid and RNA densities appear to occur near the icosahedral 3-fold axes, as evident in the equatorial section (Fig. 4B, filled white arrowhead). An additional, smaller contact appears to be located between the icosahedral 5-fold and 2-fold axes (Fig. 4B, unfilled white arrowhead).

Subunit arrangements observed in radial sections. Radial sections of the 3D reconstruction were found to complement surface views for gaining a better understanding of GLV capsid organization. Clearly distinguishable, triangular "pizza-slice" elements fully constitute the sections at higher radii (e.g., 236 to 214 Å in Fig. 4C). Visualized especially well at radius 214 Å, but also at 223 and 217 Å, is how the capsid is organized with four of these elements, representing two A and two B subunits, arrayed around each icosahedral 2-fold axis (Fig. 4C). As also inferred from the surface view, these four subunits around each icosahedral 2-fold axis appear to be arranged as two antiparallel A:B pairs, within each of which the A and B subunits make quasi-2-fold-symmetrical, tail-to-tail contacts across a broad, flat interface. These A:B contacts appear to be the major ones that connect adjacent decamers at these higher radii, although B:B contacts around the icosahedral 3-fold axes appear also to contribute. Progressing to lower radii in the sections, it is nearly impossible to distinguish individual CP subunits at radii 209 and 199 Å (Fig. 4C). Instead, at these radii almost continuous protein densities are seen, representing the core of the capsid shell. At the lowest radii in the capsid (e.g., radii 191 and 182 Å in Fig. 4C), however, the A and B subunits are again distinguishable.

The close correspondence in features between A and B subunits, across the full capsid thickness, is notable in the radial sec-

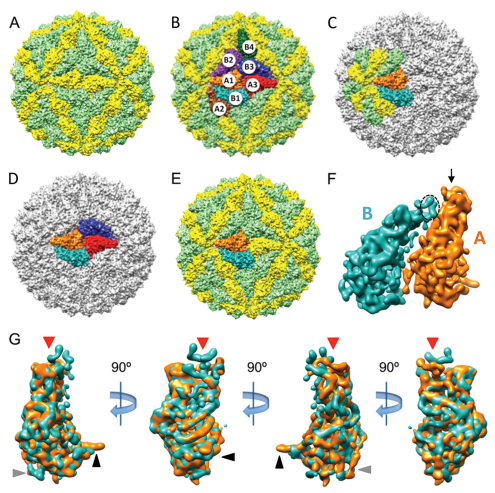


FIG 5 Segmentation of the GLV virion capsid and possible assembly intermediates. (A) Densities corresponding to CP-A and CP-B subunits according to segmentation analysis are colored yellow and light green, respectively. (B) Subunits are labeled to identify contacts between them. Colors are the same as in panel A, plus subunit  $A_1$  in orange, subunit  $B_1$  in cyan, subunit  $B_2$  in brown, subunit  $B_2$  in magenta, subunit  $B_3$  in red, subunit  $B_3$  in blue, and subunit  $B_4$  in green. (C and D) The most plausible assembly intermediates are shown: compact decamer (C) and compact tetramer (D). Colors are the same as in panel B. (E) One of the AB dimers in the GLV capsid in which the two subunits are more parallel in orientation. (F) Rotated 90° clockwise and enlarged view of the AB dimer in panel E. The black arrow indicates the region of CP-A that approaches the icosahedral 5-fold channel, and the dotted ellipse indicates the comparable region of CP-B, which has a different orientation. (G) Superimposition of segmented A and B subunits shown in four different orientations, beginning with a view as seen from outside the capsid (as in panel E) and subsequently rotated three more times, each by 90° clockwise. Predominant differences occur at the tip of each of the two subunits closest to the icosahedral 5-fold axis (red arrow) and at the AB interface near the icosahedral 2-fold axis (black/gray arrows).

tions (Fig. 4C). This correspondence pertains to not only their similar outlines but also their similar internal features appearing to represent secondary-structure elements. The A and B subunits thus appear to lie at nearly the same radial heights throughout the capsid, although close inspection suggests that the A subunits may lie just a few angstroms (i.e., 1 to 3 Å) higher. In addition, a small clockwise rotation along the longitudinal axis of the A subunits (relative to B subunits; see below) may explain subtle differences observed between the two classes of subunits, especially at lower radii (e.g., see radius 191 Å).

The positions of channels through the GLV capsid are of interest, in particular with regard to RNA transcription and RNA release from particles. Channels 10 to 15 Å wide would be large enough to permit translocation of nucleotide triphosphates and perhaps newly synthesized RNA transcripts, but not the dsRNA genome in the absence of movements of the capsid elements. In the equatorial section, open channels that span the full capsid

thickness appear to be present at the icosahedral 5-fold axes (Fig. 4B). Using the radial sections for size estimates, however, we found that this channel narrows in width to only 10 to 11 Å at radius 209 Å and remains nearly as narrow at radius 199Å (Fig. 4C). Open channels are not evident at the icosahedral 3- and 2-fold axes, although the capsid in these locations does appear to be quite thin.

Capsid segmentation. To ascribe specific features in the GLV 3D density map to the A and B subunits (Fig. 5), we used the *ab initio* approach previously used to segment the maps of HvV190S and TVV1 (38, 39). The segmentation procedure was most difficult to perform near the icosahedral 2-fold axes, where the A and B subunits from adjacent decamers seem to interact extensively in a quasi-2-fold-symmetrical arrangement (see radii 214 and 209 Å in Fig. 4C). Nonetheless, iterative refinement of density assignments, obeying the constraints described previously (39) and carried out independently by two persons, provided consistent iden-

tification of the molecular envelopes and boundaries of CP-A and CP-B (Fig. 5). Secondary-structure elements similar in shape and placement were visible as repeating units in both segmented subunits, and no evidence for major domain swapping between them was detected (Fig. 5).

Despite having different local environments, CP-A and CP-B as defined by segmentation have very similar envelopes. Superimposition of CP-A and CP-B densities yielded a correlation value of 0.91 (see Materials and Methods), which strongly suggests they share a similar tertiary structure. Indeed, the most prominent tubular density features in each subunit (ascribed to  $\alpha$ -helical elements) are nearly identically located and oriented. Each subunit has a length of maximally  $\sim 100$  Å, and as viewed radially from inside or outside the capsid, the "crust" region is about twice as wide as the "toppings" region.

CP-A and CP-B as defined by segmentation do differ at their boundaries, however. One major difference occurs at the narrow (toppings) end of each subunit. Here, CP-A contains densities that line the icosahedral 5-fold channel (Fig. 5F, arrow) and contact an adjacent, 5-fold-related A subunit (e.g., A<sub>1</sub> contacting A<sub>2</sub> in Fig. 5B) but that are not apparent at the comparable locations in CP-B. Instead, features of similar volume and shape in CP-B are oriented toward a neighboring A subunit of the asymmetric A:B pair (e.g., B<sub>1</sub> contacting A<sub>1</sub> in Fig. 5B and dotted ellipse in Fig. 5F), such that they appear to be extended outward and flipped by 90° from their position in CP-A (Fig. 5G, red arrowhead). Two other notable differences occur near the wider (crust) end of each subunit. One occurs at the outer edge of the quasi-2-fold-symmetrical A:B interaction (e.g., for A<sub>1</sub>B<sub>3</sub> in Fig. 5B and black arrowhead in Fig. 5G), on the side distal to the icosahedral 2-fold axis, where a helix-like density from CP-A extends outward toward two neighboring B subunits (e.g., A<sub>1</sub> contacting B<sub>2</sub> and B<sub>3</sub> in Fig. 5B). In contrast, this same density in CP-B does not protrude away from the subunit but instead extends in a radial direction toward the outside of the capsid. The other notable difference in this domain is a slightly shifted loop of density in CP-B near the icosahedral 3-fold axis, which contacts the other 3-fold-related CP-B subunits (e.g., B<sub>2</sub> contacting B<sub>3</sub> and B<sub>4</sub> in Fig. 5B). This same region in CP-A, though more compact compared to that in CP-B, is the one that contacts another CP-A subunit directly across the icosahedral 2-fold axis (e.g.,  $A_1$  contacting  $A_3$  in Fig. 5B).

Different responses of GLV and TVV1 virions to temperature and pH. The capacity of GLV to transmit between cells extracellularly (8, 21), whereas other known protozoal or fungal totiviruses cannot, led us to hypothesize that GLV virions may be more stable than many of these other viruses in order to withstand extracellular environments. This idea is consistent with the preceding observation that in our preparations of GLV virions (Fig. 2B and C) only a few particles appeared to have lost their dsRNA genomes after purification, in contrast to the greater number of such particles seen in preparations of TVV1 and ScV-L-A (35, 36, 39). To examine this stability difference more quantitatively, we adapted a previously described assay using RNase III to digest accessible viral genomic dsRNA (36) and used TVV1 virions for comparison with GLV virions. This assay allows for measurement of the amount of genome that becomes exposed from its protected location inside capsids upon preheating virions to different temperatures. The results showed that GLV virions were substantially more thermoresistant than TVV1 virions (Fig. 6A), a finding consistent with the greater stability of GLV.

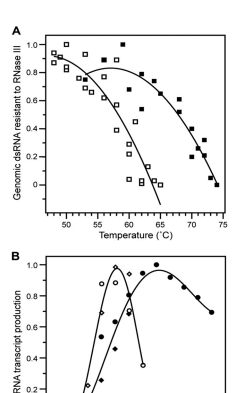


FIG 6 Different responses of GLV and TVV1 virions to temperature and pH. Experiments were performed as detailed in Materials and Methods: GLV (closed symbols), TVV1 (open symbols). (A) Normalized amount of dsRNA resistant to RNase III as a function of temperature. The data points from three (GLV) or four (TVV) experiments with each virus were superimposed in the graph and used to generate the quadratic fit. (B) RNA transcriptase activity as a function of pH. The data points from two experiments with each virus, using two different buffers (Tris-acetate [diamonds] and Tris-HCl [circles]) were superimposed in the graph and used to generate the polynomial fit.

На

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0.

Optimal growth of *T. vaginalis* trophozoites occurs at more acidic conditions (pH < 6.5) than does that of G. lamblia trophozoites (pH > 7) (54–56). Moreover, *Tritrichomonas foetus*, a relative of T. vaginalis, has been shown to possess a more acidic cytoplasm (pH 6.3) than do most eukaryotic cells, including G. lamblia (pH > 7) (55, 57). Hence, we hypothesized that the transcriptase activities of GLV and TVV1 may exhibit different pH optima, reflecting these different cytoplasmic pH values. By adapting in vitro transcriptase assays previously described for these viruses (58, 59), we found that GLV virions mediate transcription across a fairly broad pH range, peaking near pH 8.0, whereas TVV1 virions mediate transcription across a narrower range, peaking near pH 6.5 (Fig. 6B). These results are consistent with the different intracellular conditions in which these two viruses are commonly found in their two distinct hosts.

## DISCUSSION

Lack of projecting fibers in GLV. GLV appears to be more closely related than other taxonomically approved totiviruses to a group of recently discovered, nonsegmented dsRNA viruses from meta-

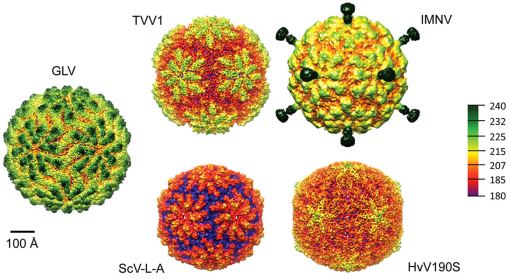


FIG 7 Comparison of GLV to four other taxonomically approved or tentatively assigned totiviruses. All particles are viewed along an icosahedral 2-fold axis, and the surfaces are color coded by radius according to the spectrum shown at right, which lists radii in angstroms. IMNV is tentatively assigned to family *Totiviridae*.

zoan hosts, including IMNV (Fig. 1). It is thus noteworthy that GLV has no projecting surface fibers, as are found at the 5-fold axes in IMNV (32) (Fig. 7). This absence of fibers is consistent with the lack of coding capacity for a fiber protein in the GLV genome, which IMNV and the related insect "totiviruses" contain upstream of their respective CPs in ORF1 (27–30). Coding for the proteolytically separated, 32-aa N-terminal peptide of the GLV CP precursor (see Fig. 3) is similarly positioned in the genome (i.e., upstream of CP in ORF1), but this peptide would be much too small to form a long fiber. It follows that in GLV, required functions for cell entry and tropism (e.g., receptor binding and membrane penetration) are likely to be mediated by as-yet-unlocalized regions of the capsid, rather than by any surface fibers as in IMNV.

In the tentatively assigned vertebrate (piscine) totivirus PMCV, a putative receptor-binding protein, containing a chemokine-superfamily motif, is encoded by a unique downstream ORF, designated ORF3 (31). This additional, recently discovered GLV-related virus (see Fig. 1) also transmits efficiently by extracellular means via virion-associated cell entry machinery. Since this property is not shared with other approved totiviruses of protozoal or fungal hosts (see Fig. 1), we suggest that GLV, IMNV and related insect viruses, and PMCV might be usefully classified in either a distinct taxonomic family or, more conservatively, in a distinct subfamily within the family *Totiviridae*.

Sequence features with potential relevance to RNA interactions and cell entry. Examination of the GLV CP precursor sequence (GenBank NC\_003555) reveals a few, previously unnoted features of interest. Immediately following the 32-aa N-terminal region that is separated by proteolysis from the CP region (10) is a strongly polar, 71-aa region dominated by charged residues (see Fig. 3). This region seems reminiscent to us of the disordered, N-terminally positioned, small R (RNA-interacting) domain of many plus-strand RNA virus CPs (60), although the R domain in those viruses is rich in basic residues, whereas this part of the sequence in GLV is acidic (calculated pI = 4.6 for this region). We hypothesize that this strongly polar region projects into the central

cavity enclosed by the GLV capsid, where its negatively charged residues may promote sliding movements of the negatively charged genomic RNA during transcription. The apparent CP contacts with RNA densities reflect where these R-domain-like regions of GLV CP are proposed to project into the interior (see Fig. 4B). These regions might also be involved in interacting with and positioning the RdRp domain of the CP/RdRp subunit(s), as has been suggested, for example, for a similar N-terminal domain in rotaviruses (61, 62).

Given the probable role of GLV CP in membrane penetration during cell entry by GLV virions into G. lamblia cells, it seemed informative to us to identify regions of CP sequence with hydrophobic character. Figure 3 highlights 14 regions of predominantly uncharged residues (regions ≥16 aa in length, including at most one charged residue), 12 of which are spread across the central portions of the CP sequence, suggestive of the folded core of this protein. In addition, there is one such region isolated near the C terminus and another near the N terminus. In fact, the N-terminal 32 aa cleaved from the CP precursor (MVWGTGYGNPSPLNPY GFASLHRGGLNPLILA) include only two charged residues (underlined), as well as a high proportion of small residues (44% Gly, Ala, Pro, and Ser), reminiscent of the membrane-penetration peptides of birnaviruses, orthoreoviruses, and others (63-66). Since only 65% of the GLV CP sequence was covered by the tryptic or chymotryptic peptides identified by LC-MS/MS analysis in this study (see Fig. 3), the possibility remains that this N-terminal peptide, or smaller peptides derived from it, might still be present in the GLV virion.

Conserved capsid elements in GLV and other *Totiviridae* members. Despite very little primary-sequence homology among the CPs of GLV, ScV-L-A, HvV190S, TVV1, and IMNV (pairwise-identity scores all <18% using EMBOSS Stretcher [http://www.ebi.ac.uk/Tools/psa/]), their capsid organizations are highly similar (Fig. 7), other than the projecting fibers in IMNV. Although the GLV capsid is somewhat larger than the others (17.5% larger than ScV-L-A; ~5–7% larger than HvV190S, TVV1, and IMNV), comparable decameric plateaus are seen on the outer surfaces of

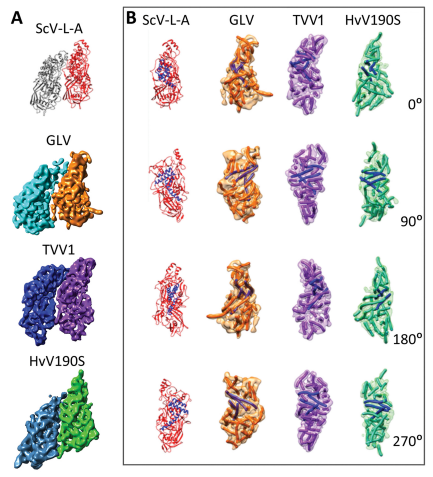


FIG 8 Comparison of the CP subunits of GLV to those of ScV-L-A, HvV190S, and TVV1. (A) Structure of the  $A_1B_1$  dimer as defined in Fig. 5. Top to bottom, as seen from outside the particle: ScV-L-A (A, red; B, gray), GLV (A, orange; B, cyan), TVV1 (A, purple; B, blue), and HvV190S (A, green; B, steel blue). The structures shown are from segmentation of cryo-TEM maps except for the X-ray crystal structures shown for ScV-L-A. (B) Structure of the A subunits of ScV-L-A, GLV, TVV1, and HvV190S, viewed from four different orientations each. The rotations indicated are around the y axis. Color scheme for the subunits is translated from panel A. Helices in the A subunit of ScV-L-A are shown as coiled ribbons, whereas putative helices in the A subunits of GLV, HvV190S, and TVV1 are depicted by tubes of similar color as their respective density maps. Four putative, long  $\alpha$ -helices, located and oriented similarly in the A subunits of these four viruses, are shown in blue.

each, although HvV190S is notably the smoothest. The larger capsid of GLV correlates with it having a larger CP than the other approved Totiviridae members shown here, although the IMNV CP is the largest (IMNV, 99 kDa > GLV, 98 kDa > HvV190S, 81 kDa > ScV-L-A, 76 kDa > TVV1, 75 kDa). The genome size of IMNV is also larger than that of GLV and the other totiviruses (IMNV, 7,561 bp > GLV, 6,277 bp > HvV190S, 5,179 bp > TVV1, 4,647 bp > ScV-L-A, 4,579 bp). Using the inner capsid diameter to estimate the central cavity volume from the density maps of these viruses, however, it turns out that GLV appears to have more space available for its genome than does IMNV (GLV  $20,035 \text{ to } 21,497 \text{ nm}^3 > \text{IMNV } 16,362 \text{ to } 19,858 \text{ nm}^3$ ; since the viral capsids are not perfectly round, the range reflects use of either the smallest or the largest inner diameter of the capsid for calculation). Assuming a single genome molecule per virion, this would result in a lower genome packing density for GLV. By being less densely packed, the GLV genome might be relatively too disorganized to give rise to concentric spherical shells of dsRNA densities in the icosahedral reconstruction as seen in IMNV and other totiviruses (32, 36, 38, 39).

Besides overall capsid organization, the individual GLV A and B subunits show a striking resemblance to those of ScV-L-A, HvV190S, and TVV1 (Fig. 8A). In addition, the orientation of GLV CP-A and CP-B with respect to each other is almost identical to the orientation of these subunits in these three viruses. The resolution of the GLV cryo-reconstruction is too low to permit a reliable trace of the entire CP polypeptide backbone, but it is clear that even at a 6.0-Å resolution the GLV CP is rich in twisted, tubular elements likely to represent  $\alpha$ -helices, as are also the CPs of ScV-L-A, HvV190S, and TVV1 (Fig. 8B). Of particular note are four putative, interacting long  $\alpha$ -helices (colored blue in Fig. 8B) that are similarly located and oriented in the CP subunits of all four viruses (best seen in the 90° and 180° side views in Fig. 8B). Thus, despite some differences in length, orientation, and angular position in the different viruses, these putative  $\alpha$ -helices appear to constitute a conserved core of the CP subunits that might be common to all totiviruses (38), including GLV.

Capsid channels and relative stability of GLV and TVV1. Following entry into the *G. lamblia* cytoplasm, the particle-enclosed GLV genome is presumably transcribed by the RdRp domain of

the CP/RdRp subunit(s) still inside the confines of the "T=2" icosahedral capsid, and the plus-strand transcripts are then released from the particle (e.g., see evidence for this in vitro activity in Fig. 6B), allowing translation of the GLV proteins and packaging into nascent, progeny GLV virions. This strategy for transcription inside the icosahedral capsid is thought to be common to all dsRNA viruses except possibly birnaviruses (67), which lack the "T=2" capsid organization (68). The only solvent-accessible channels that span the GLV capsid appear to occur at each of the 12 5-fold axes, and though somewhat narrow (ca. 10 to 11 Å in width), they are probably wide enough to permit both entry of nucleoside triphosphates and exit of newly synthesized transcripts during transcription. It is also conceivable, and perhaps likely, that conformational changes in the capsid during cell entry and/or in association with transcription may enlarge these channels or perhaps open other ones.

The 5-fold channels in GLV are considerably smaller than those in TVV1, which narrow to only  $\sim$  16 Å in the lower portions of the capsid shell (39). Although these 5-fold channels, absent conformational changes, are not wide enough in either virus for genomic dsRNA to pass through, we have found that purified preparations of TVV1 virions contain a substantial fraction of "empty" particles, i.e., particles devoid or depleted of genome, with accompanying evidence for dsRNA release following purification (39). Indeed, we have suggested that the relatively wide 5-fold channels in TVV1 may serve as the conduit for this release of genome upon limited conformational widening of these channels. This may in fact have significant biological relevance as the source of dsRNA for TLR3/TRIF-dependent signaling of proinflammatory responses by mammalian cells that encounter TVV1positive Trichomonas vaginalis parasites in the human genitourinary tract, thereby contributing to human disease (18, 19). Regarding a similar possible effect of GLV virions in contributing to human enteric disease by G. lamblia, we note that our purified preparations of GLV virions contained a much lower fraction of empty particles or evidence for dsRNA release (see Fig. 2B and C), a finding consistent with our experimental evidence that GLV virions are substantially more thermoresistant than TVV1 virions (Fig. 6A). This greater stability of GLV virions is furthermore consistent with its capacity for transmission between cells by extracellular means, entailing a greater need for it to withstand a wider range of environments but potentially reducing its propensity for proinflammatory signaling as observed for TVV1 (18, 19).

Possible pathways of GLV/totivirus capsid assembly. Assembly pathways for the capsids of icosahedral viruses often include symmetrical dimers or trimers as initial intermediates (69). Assembly of a "T=2" capsid like that of GLV, however, could not simply involve CP trimers, because only half of the subunits (60 CP-B) are arranged as 20 trimers in the capsid; the other half (60 CP-A) are arranged as 12 pentamers. Although not impossible, the formation of two such notably different intermediates by the chemically identical CP subunits seems unlikely.

"T=2" assembly pathways potentially involve compact decamers as intermediates (69), 12 of which would coalesce to form the complete capsid. Such a pathway involving compact decamers would rule out the participation of either symmetrical trimers or symmetrical dimers as intermediates and would instead rely on the formation of asymmetrical CP dimers (one A and one B subunit arranged in a near-parallel, side-by-side manner). Although this asymmetrical dimer followed by decamer assembly is cer-

tainly plausible, inspections of the GLV and other totivirus capsid structures suggest that their "T=2" capsid assembly pathway may instead involve 2-fold-symmetrical, extended CP dimers, two of which could associate to form compact, 2-fold-symmetrical CP tetramers. A related scenario is that these extended CP dimers might be transient and reversible, such that the tetramer represents the first stable intermediate. In any case, 30 such tetramers could then coalesce to form the complete capsid, possibly by first assembling into trimers of tetramers (69), 10 of which would then combine. Yet another, conceivable "T=2" assembly pathway involving a stable tetramer intermediate would have extended CP dimers continuing to accrete to an initial, nucleating tetramer.

Interestingly, studies of *Pseudomonas* bacteriophages φ6 and φ8 (family *Cystoviridae*) (70, 71), three fungal partitiviruses (family Partitiviridae) (72, 73), and rabbit picobirnavirus (family Picobirnaviridae) (65) have all identified 2-fold-symmetrical CP tetramers as the most probable assembly intermediates. Possibly, some "T=2" capsids assemble via a decamer pathway (e.g., family Reoviridae), whereas others assemble via a tetramer pathway (e.g., families Cystoviridae, Partitiviridae, and Picobirnaviridae) (69). In the case of family Totiviridae, the greater CP mass and more elongated CP structure of its members (Fig. 8) might suggest these viruses are more like members of family *Reoviridae* and assemble via the decamer pathway, but we consider the tetramer pathway to remain a viable alternative. In fact, the substantial interactions seen between CP-A and CP-B within each quasi-2-fold-symmetrical dimer in the capsid (segmentation was most difficult to perform at this subunit interface) provide direct evidence in favor of the dimer/tetramer assembly pathway.

Beyond these considerations strictly about CP subunit assembly, other key questions remain about how the CP/RdRp fusion subunit(s) and the genomic RNA assemble into mature virions. For example, regardless of the precise manner by which the GLV/totivirus CP subunits form assembly intermediates, do the one or two CP/RdRp molecules per virion always occupy either an A or a B position, or can the CP/RdRp molecule(s) assemble and function in either of these positions? Also, is the viral RNA recruited by a particular intermediate, presumably one containing a CP/RdRp subunit? These and other challenging questions need to be addressed in future studies.

### **ACKNOWLEDGMENTS**

We are grateful to C. C. Wang and coworkers (University of California, San Francisco) for providing a stock of *G. lamblia* WBI cells and to Raina N. Fichorova and coworkers (Brigham and Women's Hospital and Harvard Medical School) for providing an original stock of *T. vaginalis* UH9 cells. We also thank Norman H. Olson for technical assistance with microscopy, Maria Ericsson for negative-stain microscopy of GLV virions that had been stored at different conditions, and John M. Asara and coworkers (Beth Israel Deaconess Medical Center and Harvard Medical School) for LC-MS/MS analyses of GLV virions.

This study was supported in part by NIH grants R37 GM033050 (to T.S.B.) and R56 AI091889 (subcontract to M.L.N.). Support from the University of California San Diego and the Agouron Foundation (all to T.S.B.) was used to establish the cryo-TEM facilities used in this study.

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