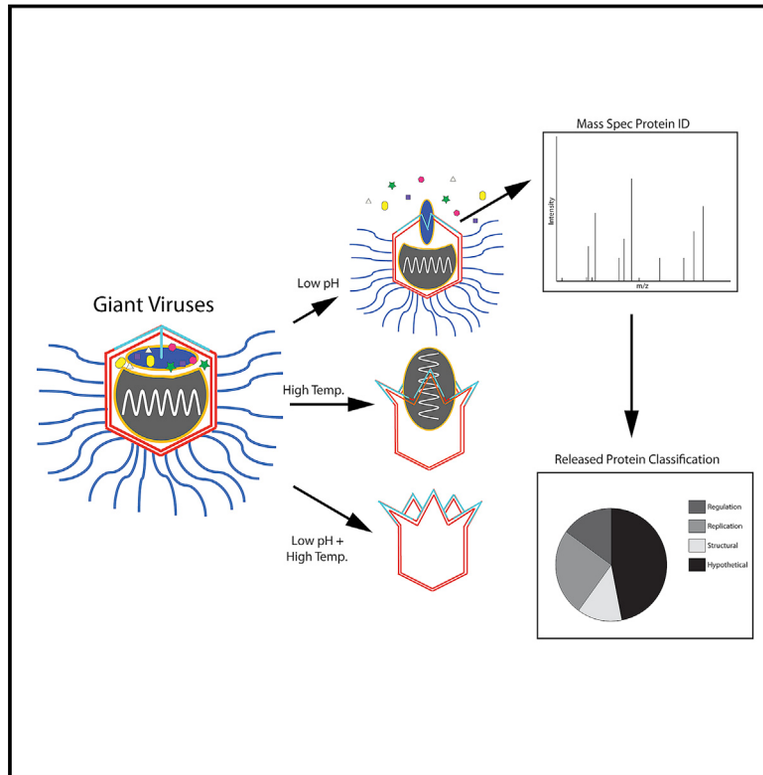


Structural and Proteomic Characterization of the Initiation of Giant Virus Infection

Graphical Abstract



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In Brief

An architectural and proteomic characterization of Mimivirus infection intermediates illustrates the conserved molecular forces driving infection and identifies proteins released from the giant virus early in the infection cycle.

Highlights

- Giant virus infection intermediates characterized by cryo-EM, cryo-ET, and SEM
- Proteomics reveal proteins released from capsids during infection
- Low pH treatment releases proteins present in the viral seed
- Low pH and high temperature fully open the stargate and release the genome

Article

Structural and Proteomic Characterization of the Initiation of Giant Virus Infection

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SUMMARY

Since their discovery, giant viruses have expanded our understanding of the principles of virology. Due to their gargantuan size and complexity, little is known about the life cycles of these viruses. To answer outstanding questions regarding giant virus infection mechanisms, we set out to determine biomolecular conditions that promote giant virus genome release. We generated four infection intermediates in Samba virus (*Mimivirus* genus, lineage A) as visualized by cryoelectron microscopy (cryo-EM), cryoelectron tomography (cryo-ET), and scanning electron microscopy (SEM). Each of these four intermediates reflects similar morphology to a stage that occurs *in vivo*. We show that these genome release stages are conserved in other mimiviruses. Finally, we identified proteins that are released from Samba and newly discovered Tupanvirus through differential mass spectrometry. Our work revealed the molecular forces that trigger infection are conserved among disparate giant viruses. This study is also the first to identify specific proteins released during the initial stages of giant virus infection.

INTRODUCTION

A hallmark of newly discovered giant viruses (GVs) is their incredibly complex biology, including gargantuan capsids and large genomes. The sheer size and complexity of these viruses, especially the inclusion of “junk” DNA in the form of introns (Azza et al., 2009; Boratto et al., 2018), challenges the canonical view of viruses as small, streamlined, and efficient killing machines. For example, most GV capsids are larger than 300 nm and many have genomes exceeding 1 MB, containing an estimated 1,000+ open reading frames (Colson et al., 2017). By contrast, some of the smallest viruses include the porcine circovirus (17 nm capsid; ~2,000 base genome; four proteins; Dhindwal et al., 2019) and the human rhinovirus (30 nm capsid; ~7,200 base genome; 11 proteins; Jacobs et al., 2013). ~69% of known viruses encode for less than 10 proteins (Brandes and Linial, 2019), highlighting the complexity of GV capsids and the true extent of our lack of knowledge concerning this new class of viruses.

GVs have been associated with a wide variety of hosts, including amoeba (Aherfi et al., 2016b), animals (Andrade et al., 2015, 2018; Dornas et al., 2014a; Khan et al., 2007), as well as human and murine cells (Ghigo et al., 2008; Lusi et al., 2017). However, amoebas also infect these creatures, casting doubt on the true GV reservoir. Although GV capsids have been associated with human respiratory diseases (Khan et al., 2007; La Scola

et al., 2005; Saadi et al., 2013a, 2013b), inflammatory conditions (Popgeorgiev et al., 2013; Shah et al., 2014), and cancers (Aherfi et al., 2016a), no direct link between GV capsids and human disease has yet been established. Despite an unusually broad host range and pathogenicity, little information is available on how GV capsids access their hosts. Infection usually occurs via phagocytosis (Abrahão et al., 2014; Ghigo et al., 2008). Once phagocytosed, a unique capsid vertex opens, which promotes nucleocapsid release and fusion with the phagosomal membrane, ultimately releasing the genome into the host cytoplasm. A pseudo-organelle (viral factory) is then formed (Suzan-Monti et al., 2007), and host replication factors are hijacked. The endpoint of GV infections is host cell death and release of new GV progeny into the environment.

GVs are ubiquitous (Aherfi et al., 2016b; Andrade et al., 2018) and maintain infectivity in harsh environments, such as alkaline lakes (Abrahão et al., 2018), frozen permafrost (Legendre et al., 2014), 3 km deep in the ocean (Abrahão et al., 2018), and Antarctic dry valleys (Andrade et al., 2018; Kerepesi and Grolmusz, 2017). GV capsids have retained infectivity following exposure to harsh chemicals (Campos et al., 2012), extreme pH and salinity (Abrahão et al., 2018), and extreme temperatures (Andrade et al., 2018; Legendre et al., 2014) and are able to persist on hospital equipment (Campos et al., 2012; Dornas et al., 2014b). To survive such extremes, GV capsids have developed incredible capsid stability. Some GV capsids can retain infectivity for 30,000 years in permafrost (Legendre et al., 2014, 2015).

Although capsid stability is beneficial for a virus to persist in harsh environments, it also creates a thermodynamic barrier that must be overcome once a suitable host cell is encountered; all known viruses must do this to propagate. Strategies and structures used for genome translocation are generally conserved across viral families. Among the tailed double-stranded DNA (dsDNA) bacteriophages (*Caudovirales*), tail complexes interact with host receptor proteins to trigger conformational changes in the virion, leading to genome release (Parent et al., 2018). Similarly, many classes of eukaryotic viruses have conserved genome release mechanisms. Most enveloped viruses, including HIV, influenza, Zika virus, and herpesvirus, utilize one of three structurally conserved membrane fusion protein varieties (Harrison, 2015). Non-enveloped viruses, such as rhinovirus, poliovirus, and adenovirus, utilize conserved capsid structures to interact with host receptors to trigger genome uncoating (Suomalainen et al., 2013).

Morphologically, GV virions are either icosahedral, exemplified by *Acanthamoeba polyphaga* mimivirus (La Scola et al., 2003), or non-icosahedral, typified by Mollivirus and Pithovirus (Legendre et al., 2014, 2015). Like their smaller cousins, GVs also share conserved capsid structures that are used during infection. In many GVs, the unique capsid vertex provides a gateway for the infection process, but they also provide a mechanism to prevent premature loss of their precious cargo. GVs have developed at least two distinct vertex structures to seal the unique vertex until the time is right for infection: “corks” and “starfish.” Non-icosahedral GVs tend to utilize one or more cork-like structures (located flush with the capsid surface) as a seal complex (Andreani et al., 2016; Legendre et al., 2014; Philippe et al., 2013). A newly discovered class of non-icosahedral GV, consisting of members such as Pandoravirus (Legendre et al., 2014) and Orpheovirus (Andreani et al., 2018), contain an ostiole-like structure, distinct from the cork-like structure.

Mimivirus-like icosahedral GVs utilize an external proteinaceous seal complex that resembles a five-pointed starfish (Klose et al., 2010; Xiao et al., 2009). These complexes sit at the outermost capsid layer at a unique 5-fold vertex (called the stargate vertex) and prevent it from opening (Xiao et al., 2009). Traditionally, both the unique capsid vertex and the external seal complex have been packaged together and called either the “stargate” or the “starfish.” We will refer to the unique capsid vertex as the stargate and the seal complex as the starfish. Non-mimivirus-like icosahedral GVs, such as PBCV-1 (Milrot et al., 2017), faustovirus (Klose et al., 2016), and Pacmanvirus (Andreani et al., 2017), do not utilize stargate vertices and have evolved alternative genome release strategies.

Starfish structures are found in diverse GVs, such as mimivirus (Klose et al., 2010; Xiao et al., 2009), Samba virus (SMBV) (Campos et al., 2014; Schrad et al., 2017), and the newly discovered tupanviruses (Abrahão et al., 2018; Silva et al., 2019), and are more common than the cork-like seals among GVs. Yet relatively little is known about the mechanism governing the stargate. The molecular forces and biochemical trigger(s), such as receptor proteins or phagosomal transitions, that facilitate stargate opening are unknown. Additionally, the fate of the starfish remains a mystery; is the complex removed from the capsid *en masse* or does the complex simply unzip?

The general steps and gross morphological changes that accompany GV infection have been visualized via thin section transmission electron microscopy (TEM) of infected cells (Abrahão et al., 2014; Silva et al., 2019). Following phagocytosis, the stargate vertex begins to open 1–3 h post-infection (Silva et al., 2019), yet little is known about the specific proteins and biomechanical forces that mediate this process. This knowledge gap is largely due to two factors: the complexity of GV virions and the lack of a robust model system for detailed biochemical and/or biophysical studies. Here, we have created an *in vitro* model system for studying the choreography that governs GV genome release. We were able to trap infection intermediates, identify specific proteins released during the initial stage of stargate opening, and test the efficacy of this technique on other icosahedral GVs, including SMBV, a member of *Mimivirus* lineage A (Campos et al., 2014); a mimivirus variant, M4 (Boyer et al., 2011); Tupanvirus soda lake (TV) (Abrahão et al., 2018); and Antarctica virus (Andrade et al., 2018). Additionally, our model reveals that members of *Mimivirus* lineage A unzip their starfish complexes to initiate infection.

RESULTS AND DISCUSSION

SMBV Is Resistant to the Vast Majority of Chemical Treatments

To probe the molecular forces that play a role in SMBV starfish complex stability, we exposed SMBV to treatments known to affect morphology in other viruses (Table S1). The effect of each treatment on particle stability was assessed via cryoelectron microscopy (cryo-EM). Treatments included the denaturants urea (≥ 9 M) and guanidinium hydrochloride (≥ 6 M), the detergent Triton X-100, organic solvents, such as chloroform and DMSO, as well as enzymes DNase I, bromelain, proteinase K, and lysozyme. None of these treatments resulted in disruption of the SMBV virion over the baseline of $\sim 5\%$ spontaneously open untreated SMBV particles (Schrad et al., 2017). Three treatments did lead to significantly increased disruption of the stargate vertex: low pH; high temperature; and high salt (see following sections). Similar treatments have been used to induce morphological changes that reflect biologically relevant structural transitions in a variety of phages and viruses (Bothner et al., 2005; Conway et al., 2001, 2007; Kant et al., 2018; Parent et al., 2010; Roos et al., 2012; Teschke et al., 2003; van de Waterbeemd et al., 2017; Yu et al., 2008; Zheng et al., 2014).

Electrostatic Interactions Are Critical for SMBV Starfish Stability

We hypothesized that pH changes occurring during and after phagocytosis may trigger SMBV stargate opening. Therefore, we dialyzed SMBV particles against different sodium phosphate buffer solutions, ranging in pH from 2 to 12 (Figure 1). Particles were visualized via cryo-EM (Figures 1A and 1B), and the percent of open particles (POP) was calculated. At and above pH 4, there was no appreciable change in the POP, compared to native (pH 7.4) levels (Figures 1A–1C). However, at and below pH 3, $\sim 60\%$ of the SMBV capsids had opened. Although the conditions that produced an increase in SMBV POP (lower than pH 4) are more acidic than the environment predicted within the amoebal

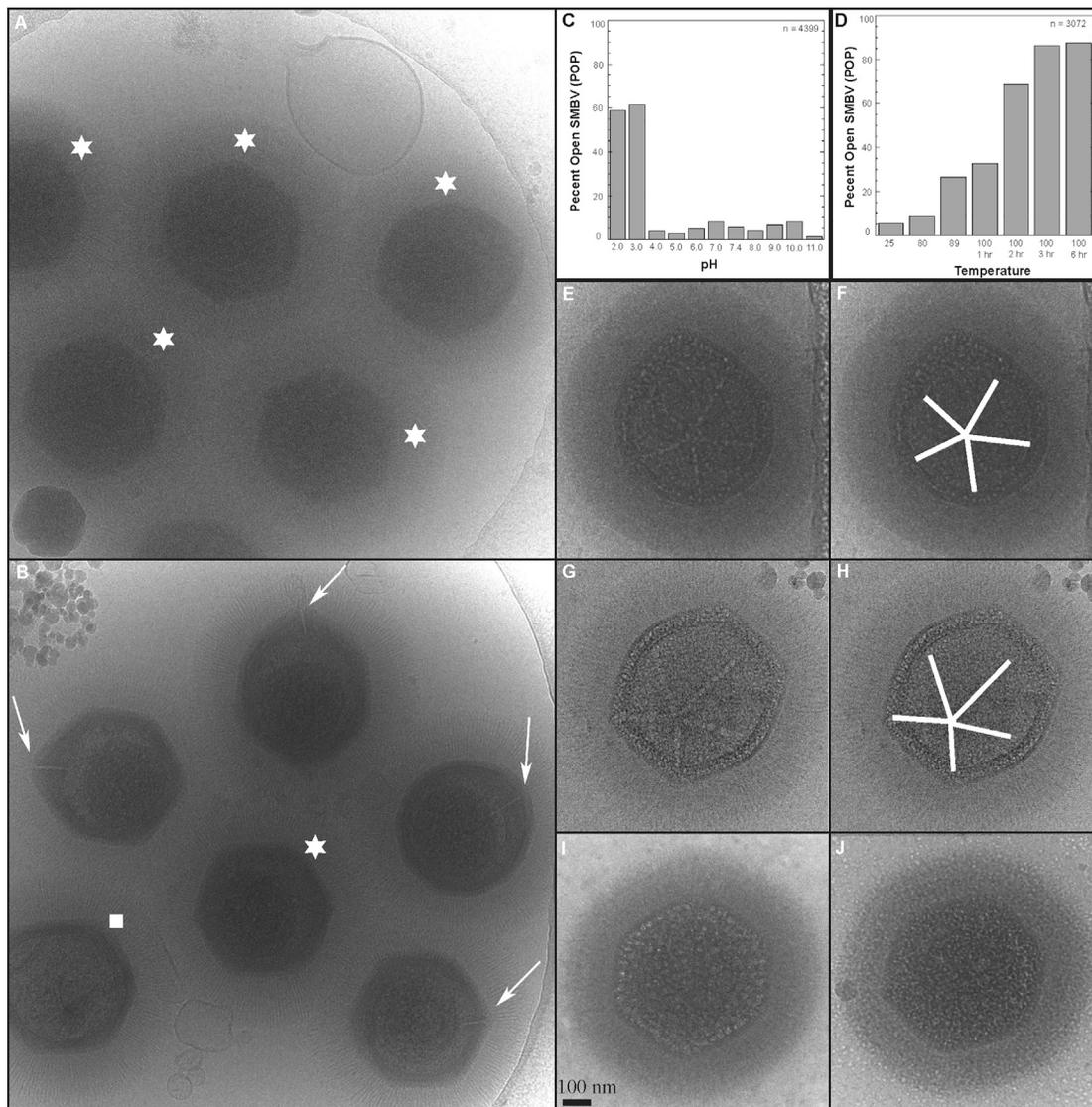


Figure 1. Low pH and High Temperature Triggered an Increase in SMBV POP

(A and B) Representative micrographs of native (A) or low-pH-treated (B) SMBV particles. Unopened particles are denoted with stars, and those with visible cracks are marked with arrows. One particle in (B), marked with a square, is clearly empty (lacks the internal density of the nucleocapsid), but cracks in the capsid are not visible, as that vertex is not contained within the micrograph. Particles such as these were counted as open when generating the percentages of open particles (POP).

(C) POP following treatment at various pH.

(D) The POP of SMBV particles incubated at elevated temperatures.

(E–H) Bubblegram images of two different native GV particles revealing star-shaped radiation damage patterns caused by the starfish complex (highlighted in F and H).

(I and J) Final micrographs in bubblegram series of two different low-pH-treated SMBV particles. Note the lack of any star-shaped radiation damage patterns. See also [Figure S1](#).

phagosome, which is \sim pH 4 (Flannagan et al., 2015; German et al., 2013; Lopez and Skaar, 2018), stargate opening is likely dependent on acidic pH, both *in vivo* and *in vitro*.

Unlike spontaneously opened GV capsids (Schrad et al., 2017; Xiao et al., 2009; Zauberan et al., 2008), these SMBV capsids were not fully open. Instead, the particles had small, noticeable cracks at one capsid vertex that assumed a star-shaped pattern. The opening of the stargate vertex at low pH is irreversible: SMBV particles returned to neutral pH still displayed star-

shaped cracks in their capsids. In some particles, the extra membrane sac was caught in the process of leaving the capsid through the newly opened vertex (Figure 2E). In other particles, the sac is not visible, suggesting that it had escaped prior to imaging. Release of the extra membrane sac, also referred to as the viral seed, has been hypothesized in other GVs. The extra membrane sac is thought to contain proteins responsible for the formation of the viral factory (Mutsafi et al., 2010; Silva et al., 2019; Suzan-Monti et al., 2007), and it is distinct from the nucleocapsid

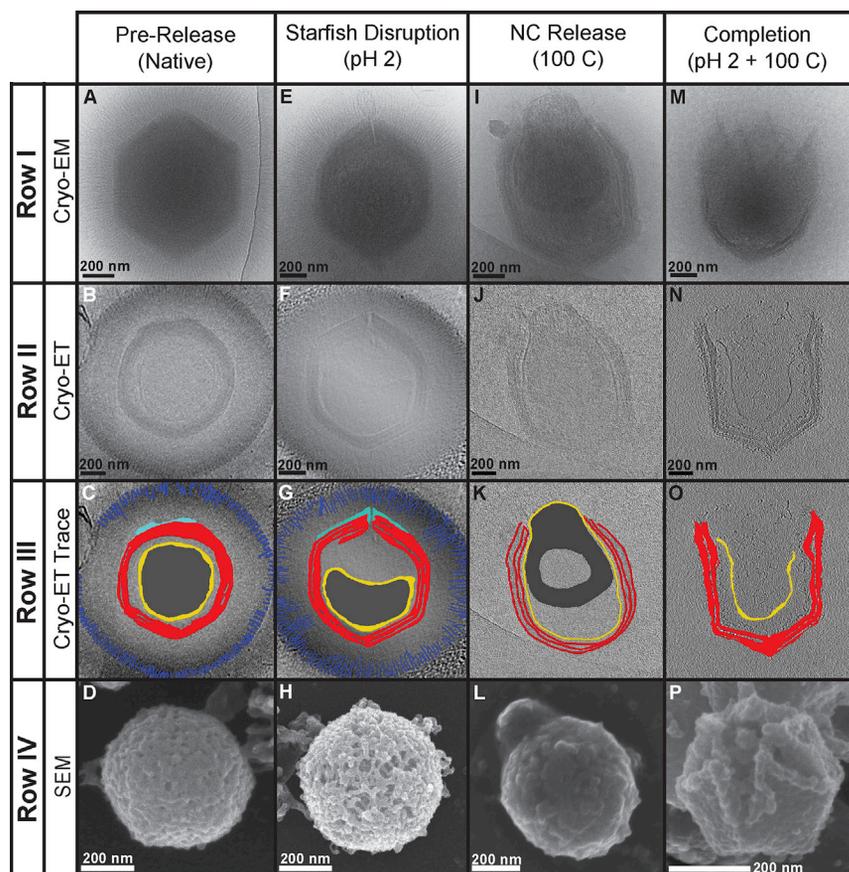


Figure 2. Electron Microscopy of SMBV Genome Release Stages

(Row I) 2D cryoelectron micrographs of particles following either no treatment (A) or post-incubation with pH 2 (E), 100°C (I), or both pH 2 + 100°C (M). (Row II) Central slices ($z = 20$) of cryoelectron tomograms of particles following either no treatment (B) or post-incubation with pH 2 (F), 100°C (J), or both pH 2 + 100°C (N). (Row III) Central slices of cryo-tomograms with key features highlighted. Blue, distal tips of the external fiber layer; cyan, starfish seal complex; dark gray, dsDNA; red, capsid; yellow, lipid membranes (nucleocapsid). Slices are shown for virions following either no treatment (C) or post-incubation with pH 2 (G), 100°C (K), or both pH 2 + 100°C (O). (Row IV) Scanning electron micrographs of particles in various stages of genome release following either no treatment (D) or post-incubation with pH 2 (H), 100°C (L), or both pH 2 + 100°C (P). See Videos S2, S3, S4, S5, S6, S7, S8, and S9 for videos of the tomograms and tilt series. See EMD-20745-20748 for tomogram volumes.

likely destabilizes through an “unzipping” mechanism rather than *en masse* release. As low pH treatment is able to trigger star-gate vertex opening *in vitro*, we conclude that electrostatic interactions play a very important role in stabilizing this vertex prior to infection. To confirm that these interactions play a role in stargate opening, we treated GV particles with 4M NaCl and

membrane when visualized in 3D using cryoelectron tomography (cryo-ET) (Schrad et al., 2017; Xiao et al., 2009). To our knowledge, this is the first study to demonstrate reproducible release of the extra membrane sac and to identify some of the proteins that may be released with this complex (below).

We could see that the particles had indeed opened following low pH treatment. Using 2D images alone, we could not, however, determine whether the starfish complex was released *en masse* or whether it remained associated with the capsid. Therefore, we used scanning electron microscopy (SEM) to probe surface features. Unfortunately, SEM images of pH-2-treated SMBV particles (Figure 2H) also did not provide definitive evidence for the presence of the starfish seal, as the layer of external fibers blocked access to the capsid surface. Enzymatic removal of the fibers was needed to visualize the surface of the starfish seal in mimivirus by SEM (Xiao et al., 2009), but unfortunately, this protocol did not remove the fibers in SMBV (Table S1). We next generated 3D reconstructions of opened SMBV particles by cryo-ET (Figure 2; Video S3; EMD-20747). Tomograms confirm that the stargate vertex, and only the stargate vertex, is open in the pH-2-treated particles. Extra density corresponding to the starfish seal is clearly observed along the edges of the outer capsid layer at the stargate vertex (Video S3). Therefore, it is likely that, at least some, if not all, of the proteins that comprise the starfish seal complex remain attached to the capsid after low pH treatment. Our tomograms suggest that the SMBV starfish

then removed the salt by dialysis for cryo-EM imaging. High salt concentrations resulted in ~40% POP (Figure S1).

We used “bubblegram” imaging, a cryo-EM technique used to localize unique features within macromolecular complexes. In this technique, samples are overexposed to produce beam-induced radiation damage. Hydrogen (H_2) gas released as a result of the radiation damaging can become trapped and sometimes produces noticeable “bubbling” in the micrograph. This bubbling can be used to reveal the location and shape of the unique features in viral capsids (Parent et al., 2018), such as phage Φ KZ inner bodies (Wu et al., 2012) and also ejection proteins in phage P22 (Wu et al., 2016).

When untreated SMBV particles were exposed to excessive electron radiation, particles produced a star-shaped radiation damage pattern (Figures 1E and 1F; Video S1). By contrast, pH-2-treated SMBV particles displayed no star-shaped pattern (Figures 1I and 1J). As expected, the lack of a star-shaped radiation damage pattern is consistent with the hypothesis that the H_2 gas is no longer being trapped in the SMBV virion, as the low pH treatment disrupted the stargate vertex seal.

Increased Thermal Energy Is Required for Nucleocapsid Release

Low pH or high salt alone was insufficient to fully open SMBV particles, indicating that electrostatic interactions are not solely responsible for fully sealing the stargate. Therefore, we next

analyzed the effect of temperature on SMBV stability. We incubated the virions for 1 h up to 100°C, assayed the virions for morphological changes using cryo-EM, and then compared these data to images of unheated particles. After 1 h at 100°C, the POP was ~33% (Figure 1D). Following an additional incubation for up to 6 h, the POP increased to a maximum of ~88%. Additional incubation at 100°C did not increase the POP further.

Unlike low pH or high salt, which simply crack the stargate vertex, high temperatures resulted in open stargate vertices and nucleocapsids in the process of exiting the virion (Figure 2; Videos S4 and S5; EMD-20748). Within these nucleocapsids, the DNA appears to have reorganized, leaving pockets of seemingly empty space (discussed below). Much of the external fiber layer is removed (Figures 2 and S1), and the extra membrane sac is also fully released from these particles. High temperature could be an alternative GV defiber method to that proposed in Kuznetsov et al. (2010), especially as this previously described technique did not defiber SMBV particles. High temperature induces a conformational change that closely mimics a structural transition that occurs during mimivirus infection as seen *in vivo* (see Figure 1, panels E and F, in Abrahão et al., 2014; Figure 6), where the nucleocapsid leaves the capsid and prepares to fuse with the amoebal phagosome membrane.

Following low-pH or high-temperature treatment, there were pockets within the SMBV nucleocapsids that appear to be devoid of DNA (Figure 2). These seemingly empty pockets are not visible in the untreated SMBV particles (Figure 2) but are seen in both low-pH- and high-temperature-treated particles. Although it is possible that the void inside of SMBV nucleocapsids could be due to the extreme conditions used, it is more likely that this is biologically relevant. These pockets are only observed in SMBV particles that have begun releasing their genome, suggesting that the DNA may undergo reorganization during this process. The SMBV genome contains various chromosome condensation and histone-like proteins that could be used for this function, although their exact role in GV biology has not yet been experimentally determined. Mass spectrometry experiments (described below and shown in Table 1) suggest that many of these proteins remain with the nucleocapsid after the initial opening stage. Genome reorganization is an important stage of many virus infection processes, including HIV (Freed, 2015) and adenovirus (Mangel and San Martín, 2014). We hypothesize that genome rearrangement is also important for facilitating GV genome release into the host.

A Combination of Low pH and High Temperature Results in Complete SMBV Genome Release

Individually, low pH and high temperature had different physical effects on SMBV. These disparate treatments are affecting both electrostatic interactions and the availability of thermal energy, respectively, and each appears to contribute to SMBV virion stability. Therefore, we hypothesized that combining low pH and high temperature might have a compound effect on stargate opening. Again, following treatment, the SMBV particle morphology was analyzed via cryo-EM (Figure 2M), cryo-ET (Figures 2N and 2O), and SEM (Figure 2P). These particles have completed the entire genome release process, as seen by the absence of the nucleocapsid. Additionally, SMBV particles

were completely defibered and the internal capsid layer(s) appeared to be less rigid than the outer capsid layer (Figure 2O; Videos S6, S7, S8, and S9; EMD-20745 and EMD-20746).

SEM of dual-treated SMBV particles (Figure 2P) provides further evidence for the starfish seal. Particles treated with both low pH and high temperature clearly contain extra density around the edges of the stargate vertex, corresponding to the starfish seal. This extra density is consistent with our cryo-ET data described above, where, rather than completely dissociating from the capsid *en masse*, the starfish seal unzips while still retaining contacts with the capsid.

Molecular Forces that Stabilize the SMBV Stargate Vertex Are Conserved among Mimiviruses

We tested combining pH and high temperature on three other GVs: Antarctica virus (Andrade et al., 2018); Tupanvirus (Abrahão et al., 2018); and mimivirus M4 (Boyer et al., 2011). Although all four viruses analyzed here are mimivirus-like icosahedral GVs, these viruses encompass two separate GV clades belonging to the *Mimiviridae* family: of the genus *Mimivirus* (SMBV, M4, and Antarctica) and the proposed genus *Tupanvirus* (TV; Rodrigues et al., 2019b). Note that Tupanvirus is distinct from the others, as it has a long tail attached to the icosahedral head as seen in Abrahão et al. (2018) and Rodrigues et al. (2019a, 2019b). Following treatment, each virus was characterized via SEM and cryo-EM (Figures 3 and 6). All four GVs tested in this study opened their stargate vertices and released their nucleocapsids after being boiled in acid. Two also appeared to lose the majority of their fibers during treatment, with Tupanvirus being the exception. The fate of the fibers and their role during infection is not well known, although it has been suggested that fibers aid in viral attachment and increase phagocytosis (Rodrigues et al., 2015). These data strongly indicate that the general forces that stabilize virions and facilitate infection are conserved among distantly related amoeba-infecting members of *Mimiviridae*.

Although the general forces appear to be highly conserved, some specific mechanisms of starfish disruption are likely conserved only within distinct lineages. In our SEM data, Antarctica and mimivirus particles (Figures 3A and 3D, respectively) displayed density along the edges of the open stargate vertices, similar to the density seen in SMBV (Figures 2P and 3C). The presence of this extra density suggests that, like SMBV, the Antarctica and mimivirus starfish complexes unzip to facilitate stargate opening and genome release. TV, on the other hand, does not display this extra density (Figure 3B), suggesting that the TV starfish may completely dissociate from the capsid *en masse* during infection. TV particles also appear to fully open their stargate vertices following low-pH treatment alone. In total, our data suggest that the mechanism of seal complex unzipping may be conserved among *Mimiviridae*, with slight deviations present between the *Mimiviruses* and the proposed *Tupanvirus* genus.

GVs have changed our canonical view of virology, defying the previously known limits of capsid sizes and stabilities. The description of a new GV genome release strategy signifies another paradigm shift in our understanding of virology. As mentioned previously, smaller viruses tend to share conserved genome release mechanisms. This conservation can be

Table 1. Identification of Proteins Released from SMBV and TV Capsids

Protein	ID	Category	Presence		Ratio of Ratios	
			Band	Shotgun	Up in Supe	Down in Pellet
Samba Virus						
Actin ^{a,b}	GenBank: CAA23399.1	S	+	+	+	
Rpl7A, partial ^a	GenBank: AAY21190.1	–		+	+	
Amine oxidase	GenBank: AHJ39955.1	M		+		+
DNA-dependent RNP subunit RPB9	GenBank: AHJ39967.2	TI		+		
Ubiquitin-conjugating enzyme e2	GenBank: AHJ39993.2	M	+	+		
WD repeat-containing protein	GenBank: AHJ40002.1	–		+		
B-type lectin protein	GenBank: AHJ40019.2	S		+		
Protein phosphatase 2c	GenBank: AHJ40032.1	Rg	+			
Formamidopyrimidine-DNA glycosylase	GenBank: AHJ40038.1	Ho	+			
Hypothetical protein	GenBank: AHJ40051.1	H/TM	+			
Poly(A) polymerase catalytic subunit	GenBank: AHJ40056.1	Tx	+			
Hypothetical protein	GenBank: AHJ40060.1	H		+		+
Hypothetical protein	GenBank: AHJ40061.1	H		+		+
Thioredoxin domain-containing protein	GenBank: AHJ40071.1	Ho	+			
mRNA-capping enzyme	GenBank: AHJ40083.1	Tx		+		+
Putative FtsJ-like methyltransferase	GenBank: AHJ40084.1	Tx	+			
Hypothetical protein	GenBank: AHJ40087.2	H	+			
Low-complexity protein	GenBank: AHJ40093.1	H		+		
Core protein	GenBank: AHJ40101.1	S	+			
Hypothetical protein	GenBank: AHJ40107.2	H		+		
Capsid protein 1	GenBank: AHJ40114.2	S	+			+
Hypothetical protein	GenBank: AHJ40128.1	H	+			
Thioredoxin domain-containing protein	GenBank: AHJ40129.2	Ho	+	+	+	+
Hypothetical protein	GenBank: AHJ40139.1	H		+	+	
Hypothetical protein	GenBank: AHJ40144.1	H		+		
DNA-directed RNA polymerase subunit I	GenBank: AHJ40151.2	TI	+			
Hypothetical protein	GenBank: AHJ40159.1	H	+			
Hypothetical protein	GenBank: AHJ40160.2	H		+		
Hypothetical protein	GenBank: AHJ40162.1	H	+			
Hypothetical protein	GenBank: AHJ40169.1	H		+		
DNA-directed RNAP subunit 1	GenBank: AHJ40172.1	TI		+		+
Hypothetical protein	GenBank: AHJ40183.2	H		+	+	
Alpha beta hydrolase/esterase/lipase ^b	GenBank: AHJ40190.1	I/E	+			
Hypothetical protein	GenBank: AHJ40207.1	H	+			
Hypothetical protein	GenBank: AHJ40211.1	H	+			
Hypothetical protein	GenBank: AHJ40213.2	H	+	+	+	+
Hypothetical protein	GenBank: AHJ40220.1	H	+			
Hypothetical protein	GenBank: AHJ40230.1	H		+		
Hypothetical protein	GenBank: AHJ40243.1	H		+		+
Mannose-6P isomerase	GenBank: AHJ40247.1	M		+	+	
Hypothetical protein	GenBank: AHJ40254.1	H		+		
Hypothetical protein	GenBank: AHJ40271.2	H		+		
Tat pathway signal sequence domain protein ^b	GenBank: AHJ40276.1	I	+		+	
Collagen-like protein 7	GenBank: AHJ40290.2	S		+	+	+
Hypothetical protein	GenBank: AHJ40316.2	H		+		+

(Continued on next page)

Table 1. Continued

Protein	ID	Category	Presence		Ratio of Ratios		
			Band	Shotgun	Up in Supe	Down in Pellet	
Hypothetical protein	GenBank: AHJ40318.2	H		+			
Hypothetical protein	GenBank: AHJ40319.1	H	+				
Hypothetical protein	GenBank: AHJ40326.2	H		+	+		
Low-complexity protein	GenBank: AHJ40329.1	H	+		+		
Hypothetical protein	GenBank: AHJ40333.1	H/TM		+	+		
Chemotaxis protein	GenBank: AHJ40337.1	I/S	+				
Hypothetical protein	GenBank: AHJ40339.1	H	+			+	
Hypothetical protein	GenBank: AHJ40340.1	H	+				
Ubiquitin thioesterase	GenBank: AHJ40341.2	M	+				
Hypothetical protein	GenBank: AHJ40367.2	H/TM	+				
Virion-associated membrane protein	GenBank: AHJ40371.2	I	+				
Lanosterol 14-alpha-demethylase	GenBank: AHJ40393.1	I		+		+	
Hypothetical protein	GenBank: AHJ40423.1	H	+		+		
Collagen triple helix repeat containing protein	GenBank: AMK61745.1	S		+		+	
Choline dehydrogenase-like protein	GenBank: AMK61776.1	M	+			+	
DNA topoisomerase 1b	GenBank: AMK61799.1	Rg	+				
Probable glutaredoxin	GenBank: AMK61800.1	Rp/Ho	+			+	
Hypothetical protein	GenBank: AMK61829.1	H	+	+	+		
Hypothetical protein	GenBank: AMK61837.1	H		+		+	
Hypothetical protein	GenBank: AMK61849.1	H/TM		+			
Hypothetical protein	GenBank: AMK61856.1	H	+			+	
Regulator of chromosome condensation ^c	GenBank: AMK61866.1	Rg/I	+			+	
Thiol protease	GenBank: AMK61869.1	E	+				
Hypothetical protein	GenBank: AMK61892.1	H		+			
Hypothetical protein	GenBank: AMK61902.1	H	+				
Anaerobic nitric oxide reductase transcription regulator NorR	GenBank: AMK61903.1	Rg	+				
Ankyrin repeat protein	GenBank: AMK61918.1	–	+				
Hypothetical protein	GenBank: AMK61920.1	H	+				
Hypothetical protein	GenBank: AMK61935.1	H	+				
Hypothetical protein	GenBank: AMK61942.1	H		+		+	
N-acetyltransferase	GenBank: AMK61955.1	M	+				
Prolyl 4-hydroxylase	GenBank: AMK61959.1	Ho		+			
Proline rich protein	GenBank: AMK61968.1	–	+			+	
Hypothetical protein	GenBank: AMK61977.1	H	+	+			
NHL repeat-containing protein	GenBank: AMK61987.1	–		+			
Hypothetical protein	GenBank: AMK62013.1	H		+			
Hypothetical protein	GenBank: AMK62059.1	H	+			+	
Hypothetical protein	GenBank: AMK62082.1	H	+				
Choline dehydrogenase-like protein	GenBank: AMK62096.1	M		+		+	
Ubiquitin ^{a,d}	GenBank: CAA53293.1	M		+		+	
Tupanvirus Soda Lake							
Hypothetical protein	GenBank: AUL78681.1	E/H	+	+	+	+	
Hypothetical protein	GenBank: AUL77600.1	E/H		+			
Putative ORFan	GenBank: AUL77729.1	H		+			
Putative ORFan	GenBank: AUL78088.1	H	+	+	+	+	

(Continued on next page)

Table 1. Continued

Protein	ID	Category	Presence		Ratio of Ratios	
			Band	Shotgun	Up in Supe	Down in Pellet
Hypothetical protein	GenBank: AUL78481.1	H		+		
Hypothetical protein	GenBank: AUL78232.1	H/Rg		+		+
Hypothetical protein	GenBank: AUL78466.1	H	+	+	+	+
Hypothetical protein	GenBank: AUL77936.1	H		+		
Hypothetical protein	GenBank: AUL78214.1	H		+		+
Hypothetical protein	GenBank: AUL77907.1	H	+	+	+	+
Hypothetical protein	GenBank: AUL78468.1	H	+	+	+	+
Hypothetical protein	GenBank: AUL77723.1	H	+	+	+	+
Hypothetical protein	GenBank: AUL78464.1	H	+			
Hypothetical protein	GenBank: AUL78055.1	H		+	+	+
Hypothetical protein	GenBank: AUL77930.1	H		+	+	+
Putative ORFan	GenBank: AUL78635.1	H	+			+
Hypothetical protein	GenBank: AUL77752.1	H		+		
Hypothetical protein	GenBank: AUL78219.1	H		+		
Hypothetical protein	GenBank: AUL78093.1	H	+			+
Hypothetical protein	GenBank: AUL78067.1	H		+		
Hypothetical protein	GenBank: AUL78191.1	H	+			+
Hypothetical protein	GenBank: AUL78287.1	H		+		+
Hypothetical protein	GenBank: AUL77694.1	H	+			
Hypothetical protein	GenBank: AUL77820.1	H	+			
Hypothetical protein	GenBank: AUL78135.1	H		+		+
Hypothetical protein	GenBank: AUL78143.1	H	+			+
Hypothetical protein ^c	GenBank: AUL78348.1	H/Rg	+		+	+
Hypothetical protein	GenBank: AUL77688.1	H/TM	+			+
Hypothetical protein	GenBank: AUL78288.1	H/TM		+		+
Hypothetical protein	GenBank: AUL77718.1	H/TM/E	+			
Cu-Zn superoxide dismutase ^e	GenBank: AUL78503.1	Ho	+			
mg709 protein ^e	GenBank: AUL77661.1	Ho		+		+
Thioredoxin domain-containing protein	GenBank: AUL77963.1	Ho		+		+
Catalase HP11	GenBank: AUL78097.1	Ho		+		+
Ig family protein	GenBank: AUL78630.1	I	+			
Phosphatidylethanolamine-binding protein-like protein	GenBank: AUL77474.1	I	+			
Putative N-acetyl transferase	GenBank: AUL77680.1	M		+		
Arylsulfatase	GenBank: AUL78269.1	M		+		
Ubiquitin domain-containing protein ^d	GenBank: AUL78040.1	M	+	+	+	+
Glyoxalase	GenBank: AUL78134.1	M	+			
Putative protein kinase	GenBank: AUL78629.1	Rg		+		+
Glutaredoxin	GenBank: AUL78724.1	Rp/Ho		+	+	+
SNF2 family helicase	GenBank: AUL77941.1	Rp/Rg	+			
Capsid protein 1	GenBank: AUL78147.1	S	+			
Putative fibril associated protein	GenBank: AUL78400.1	S		+		
Kinesin-like protein ^a	GenBank: AUL77838.1	S		+		+
Major core protein	GenBank: AUL78082.1	S	+			
Putative pore coat assembly factor	GenBank: AUL78211.1	S		+		+
Mimivirus elongation factor aef-2	GenBank: AUL78714.1	TI	+			
DNA-directed RNAP subunit	GenBank: AUL78016.1	TI		+	+	

(Continued on next page)

Table 1. Continued

Protein	ID	Category	Presence		Ratio of Ratios	
			Band	Shotgun	Up in Supe	Down in Pellet
Intein-containing DNA-directed RNAP subunit 2	GenBank: AUL78362.1	TI		+		
DNA-directed RNAP subunit 6	GenBank: AUL78368.1	TI		+	+	+
DNA-directed RNAP subunit 1	GenBank: AUL78302.1	TI		+		
Putative ATP-dependent RNA helicase	GenBank: AUL77829.1	Tx/TI	+			
Actin ^{a,b}	GenBank: CAA23399.1	S		+	+	

^a*Acanthamoeba castellanii* proteins.

^bProteins similar to *Iridovirus* UPP-associated proteins.

^cProteins involved in genome rearrangement.

^dProteins directly involved in a putative ubiquitin-proteasome degradation pathway.

^eMetal-conjugating proteins.

observed within viral families, such as *Flaviviridae* (fusion proteins; Apellániz et al., 2014), *Caudovirales* (tail complexes; Parent et al., 2018), or *Orthomyxoviridae* and *Paramyxoviridae* glycoproteins (Kordyukova, 2017). This conservation also occurs across viral kingdoms. The herpesvirus portal complex shares structural similarity with many bacteriophage portal proteins (McElwee et al., 2018; Newcomb et al., 2001), and the adenovirus spike protein is homologous with the bacteriophage Sf6 tail needle knob protein (Parent et al., 2012). Mimivirus-like GVs have eschewed these known genome release structures and appear to have forged their own mechanisms, as exemplified by the common stargate mechanism.

Numerous Proteins Are Released from GV Capsids during Stargate Opening

As obvious morphological changes occurred in GVs during low-pH and high-temperature treatments, we hypothesized that proteins were likely released from the capsids at each of these stages. We analyzed proteins that remained within the SMBV and TV capsids and proteins liberated from the capsids after each treatment. We used four conditions: native virions (pH 7.4; 25°C); low pH (pH 2; 25°C); high temperature (pH 7; 100°C); and a combination (pH 2; 100°C). We then performed pellet/supernatant separations to physically separate the virions from released proteins and analyzed the contents of each via SDS-PAGE (Figure 4). A preparation scheme can be seen in Figure S2. Antarctica and mimivirus both showed a similar banding pattern as SMBV. We did not perform mass spectrometry (MS) experiments with these viruses, as there is no annotated Antarctica virus genome and mimivirus and SMBV are highly similar (Campos et al., 2012).

For both SMBV and TV, distinct proteins were released from the capsid following low-pH treatment. Some of these proteins align with proteins in the native capsid (pellet) lane, suggesting they had been released from the capsid without significant modification/cleavage. Other proteins, especially in the TV sample, did not match proteins in the native capsid lane. These bands likely represent proteins that were cleaved during treatment. For both viruses, the native supernatant lanes did not contain any visible protein bands. When the particles were incubated at 100°C (with or without prior pH 2 treatment), it appeared

that the majority of proteins were proteolytically cleaved and ran as a continuous smear on the gel, preventing detailed analysis of these samples.

Identifying the Proteins Released from SMBV and TV Virions at the Initiation of Infection

To characterize the proteins released during the initial stages of GV infection, we used MS. Initially, we used in-gel digestion of bands from the pH-2-treated SMBV and TV supernatant samples. Trypsinized fragments were analyzed via liquid chromatography-tandem mass spectrometry (LC-MS/MS), and the resultant peptides were compared to published SMBV and TV genome sequences (GenBank: KF959826.2 and GenBank: KY523104.1) as well as the *A. castellanii* genome (GenBank: KB007974.1) to identify host proteins. The *A. castellanii* actin protein was retained within these results, as this protein is known to play a role in the infection and genome release processes of *Iridoviruses* (Huang et al., 2018). From these initial data, we identified 48 SMBV and 26 TV proteins that are released from the virion following low-pH treatment (labeled with a [+] in the “band” column of Table 1).

Excising visible gel bands for MS analysis has the potential to miss proteins within the sample: some bands may be too faint to detect, some proteins may be too large or too small to be fully resolved or extracted, etc. Therefore, we also analyzed SMBV and TV samples using shotgun proteomics to maximize coverage in our study. We analyzed low-pH pellet and supernatant samples, as well as the untreated virus using the scheme shown in Figure S2. From this experiment, we identified 43 SMBV proteins and 37 TV proteins ([+] in the “shotgun” column of Tables 1 and S3). Of these proteins, 5 SMBV proteins and 7 TV proteins were previously identified from analysis of the gel bands.

In total, 78 SMBV proteins and 61 TV proteins were identified as having been released from the capsids at low pH. TV was isolated from an environment with high salinity and alkaline pH (9–12; Abrahão et al., 2018). SMBV, on the other hand, was isolated from a tributary of the Amazon River, a relatively neutral environment. Due to its location, TV had to evolve pH stability into its capsid to a greater extent than SMBV. Although TV was originally isolated from a basic environment, some of the strategies that

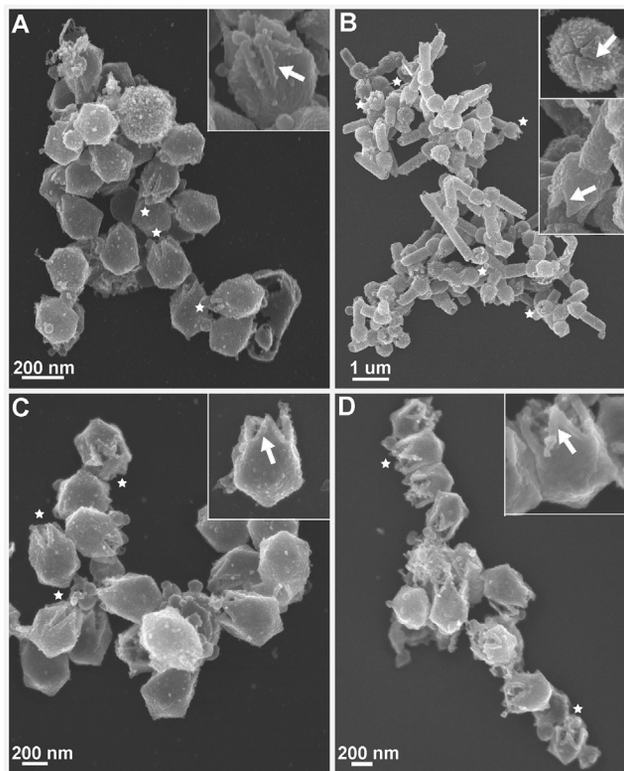


Figure 3. Post-genome Release Particles from Four GVs

Scanning electron micrographs of low-pH- and high-temperature-treated (A) SMBV, (B) TV, (C) Antarctica virus, and (D) mimivirus particles. Insets demonstrate enlarged views highlighting capsids where clear retention of the starfish seal can be seen in SMBV, mimivirus, and Antarctica particles. Asterisks in the main panels depict selected particles with clearly visible open stargate vertices.

the virus could have developed to stabilize its proteins, such as using a higher percentage of non-polar amino acids, could also stabilize the proteins at low pH.

187 and 169 total proteins were identified within the untreated mature virions of TV and SMBV, respectively (Figure 5). To identify proteins that had been released, we calculated the percent of the total peptide signal for each. We compared these percentages across the three samples, specifically looking at the ratios of supe:MA (material applied) and pellet:MA. Proteins where the supe:MA > 1 was enriched in the treated supernatant sample, indicating that they had been released from the capsids. These proteins are identified with a (+) in the “up in supe” column of Table 1. Conversely, proteins with pellet:MA < 1 were less abundant in the treated pellet than the native particles and likely also released. These proteins are identified with a (+) in the “down in pellet” column of Table 1. Proteins that are enriched in the supernatant samples are definitely released from the GV capsids, as no proteins were identified in the untreated supernatant samples (data not shown). Proteins that are depleted in the pellet samples are also likely released from the GV particles, although it is unlikely that any of these proteins are completely absent from the pellet samples (see POP in Figure 1A).

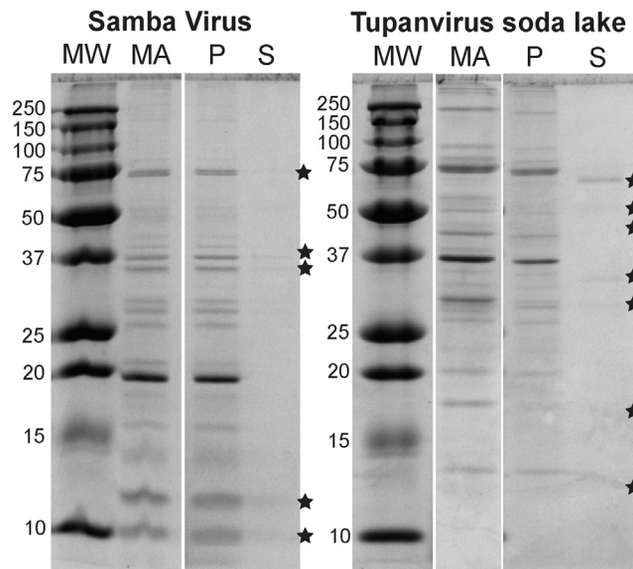


Figure 4. SDS-PAGE of pH-2-Treated SMBV and TV

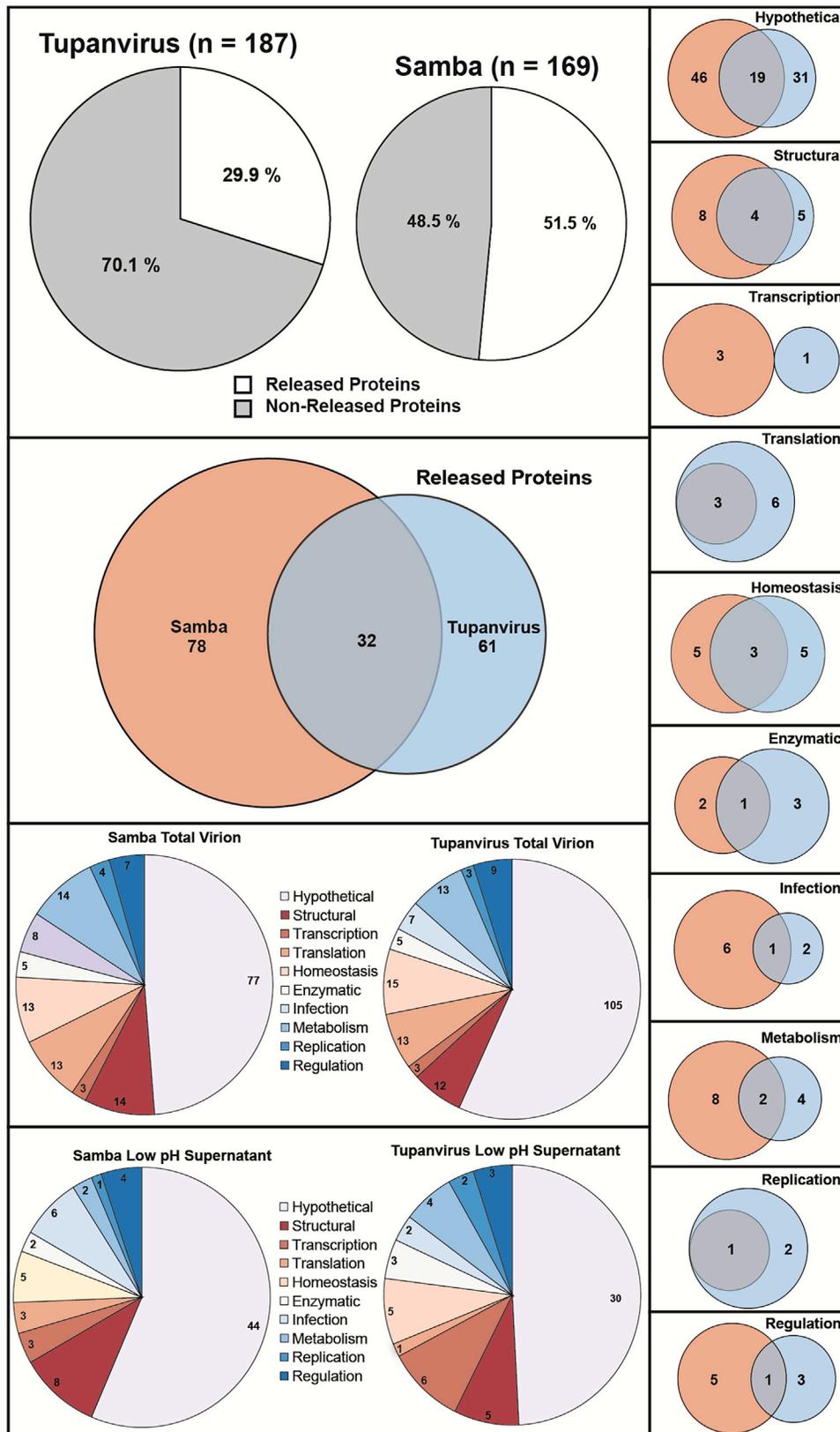
SDS-PAGE of SMBV and TV. MA, material applied (untreated viral particles); p, pellets from pH-2-treated virions; S, supernatants from pH-2-treated virions. Visible bands of proteins released into the supernatant are highlighted with asterisks. See Figure S2 for the sample preparation scheme.

SMBV releases a higher number and percentage of these proteins (86; 51.5%) than TV particles (56; 29.9%). Putative functions for the released proteins were determined via (1) previous annotation (Abrahão et al., 2018; Campos et al., 2014), (2) NCBI BLAST analysis, (3) HHBLITS analysis (Remmert et al., 2011), (4) InterPro functional prediction (Mitchell et al., 2019), and (5) PSIPRED domain prediction using DomPred (Buchan and Jones, 2019; Jones and Cozzetto, 2015). Released proteins for each virus were separated into the following 10 categories: hypothetical; structural; transcription; translation; homeostasis; enzymatic; infection; metabolism; replication; and regulation (Figures 5B–5N).

Many of the proteins released for each virus (more than 50% for SMBV) are hypothetical proteins or proteins with unknown function. 19 of the hypothetical proteins released by the two viruses displayed obvious homology between SMBV and TV (BLAST results or functional homology prediction). All of the released SMBV proteins predicted to be involved in both translation and replication had homologs among the released TV proteins. The proteins predicted to be involved in transcription and regulation, on the other hand, did not show any readily apparent homology. The homology between the released TV and SMBV proteins in general and within each category can be found in Figure 5.

Expected Protein Types Are Released from SMBV and TV Virions during Genome Release

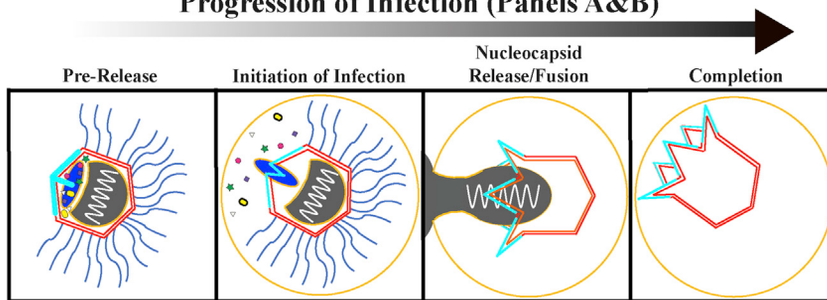
GVs need to carry out the same basic stages of the viral life cycle as their smaller cousins to replicate. Common stages include genome translocation into the host cell, blocking host replication, hijacking host machinery to make viral proteins, and making



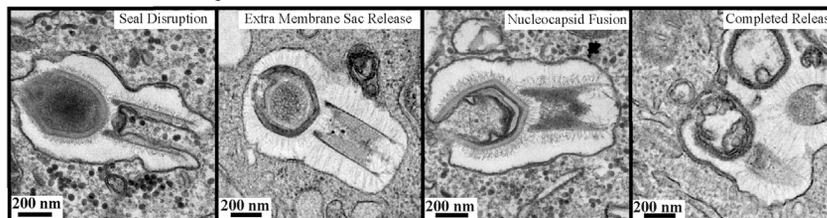
(legend on next page)

A Cartoon Model of GV Infection

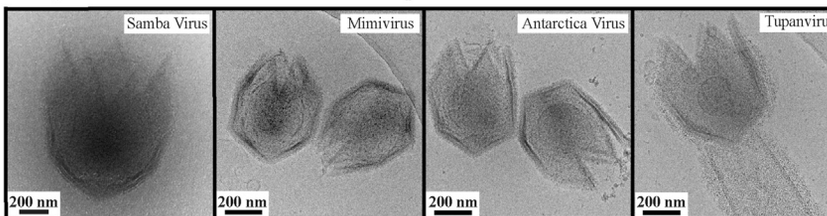
Progression of Infection (Panels A&B)



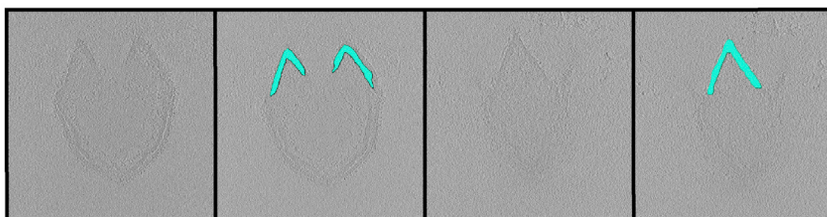
B Thin Sections of Tupanvirus Infection *In Vivo*



C Cryo-EM of GV Genome Release Completion *In Vitro*



D Slices of an SMBV Tomogram Highlighting the External Seal Complex



new viral proteins. Both SMBV and TV likely release proteins that are predicted to perform these functions, as many smaller viruses release whole proteins or peptides to facilitate this function (Manning et al., 2018; Parent et al., 2018; Wu et al., 2016). Hypothetical or unknown function proteins released from GV particles likely aid in performing these critical functions, as many of them are released during the initial phase of opening. Aside from identifying putative functions for some hypothetical proteins discussed below, determining the specific function of all of these proteins lies beyond the scope of this study.

Before the virus is able to hijack the host machinery and begin replication, it must enter the cell and translocate its genome across the phagosomal membrane into the cytoplasm. SMBV

Figure 6. Giant Virus Genome Release Process

(A and B) The progression of GV genome release depicted as a cartoon (A) and through thin section TEM images (B). The thin-section micrographs represent Tupanvirus particles visualized at varying stages of the infection process *in vivo*. The panels for “seal disruption” and “nucleocapsid fusion” were previously published in [Abranches et al. \(2018\)](#) and shown under Creative Commons Attribution 4.0 International License, which permits unrestricted use, distribution, modification, and reproduction. The remaining two panels are original images.

(C) Cryoelectron micrographs of the completion of the *in vitro* genome release process for the four GVs studied here.

(D) Z slices of an SMBV tomogram of fully opened particles highlighting the density corresponding to the starfish seal.

releases putative membrane proteins, such as a virion-associated membrane protein (AHJ40731.1) and as well as hypothetical proteins with predicted transmembrane domains that may play a role in membrane fusion (“H/TM” in [Table 1](#)). Both the extra membrane sac and the nucleocapsid must fuse with the phagosome to deliver the viral seed and dsDNA genome into the cytoplasm. This fusion event can clearly be seen in micrographs of infected cells ([Figure 6](#)). We will investigate the fusion capability of these newly identified proteins directly in future studies. Therefore, the results of this study help to assign putative roles to many proteins with previously unknown function, highlighting the power of this new method.

Additionally, both SMBV and TV release proteins were predicted to play a role in an ubiquitin-proteasome degradation pathway (UPP) (delineated by ^d in [Table 1](#)). These proteins are known to facilitate genome release in other viruses, including the large, but not quite giant, Iridoviruses ([Huang et al., 2018](#)) and herpesviruses ([Greene et al., 2012](#)). In Iridovirus infection, the UPP is coupled with metabolic, cytoskeletal, macromolecule biosynthesis, and signal transduction proteins to facilitate infection ([Huang et al., 2018](#)). Proteins predicted to carry out these functions are released from both the SMBV and TV virions alongside the UPP-related proteins (^p in [Table 1](#)). Hypothetical proteins with additional functional information predicted via BLAST, HHBLIST, PSIPRED, or InterPro are listed in [Table S2](#).

Figure 5. Comparison of Proteins Released by SMBV and TV Soda Lake

Venn diagrams comparing the total protein content and proteins released following low pH treatment of SMBV and TV particles. The homology present within these protein sets is depicted. See [Table S2](#) for hypothetical proteins with predicted transmembrane domains and [Table S3](#) for the relative abundance of individual proteins. See also [Figure S2](#).

Following genome translocation, the virus forces the cell machinery to transition from making new cellular products to making viral components. Both SMBV and TV release various subunits of a DNA-dependent RNA polymerase (SMBV: AHJ39967.2, AHJ40151.2, and AHJ40172.1; TV: AUL78016.1, AUL78362.1, AUL78368.1, and AUL78302.1). This series of proteins is critical for the life cycle of the virus, as it directs the cellular machinery of the host to recognize viral DNA in lieu of cellular DNA. These proteins, especially the various DNA-dependent RNA polymerases, may play a role in transcription as hypothesized to occur following stargate opening but before nucleocapsid release (Mutsafi et al., 2014). Additional proteins in this category likely include some of the metabolic proteins released by the viruses, especially the catabolic proteins that may play a role in degrading host defenses and machinery. These proteins include a SMBV thiol protease (AMK61869.1), a SMBV amine oxidase (AHJ39955.1), and a hypothetical TV protein with a predicted inosine/uridine-favoring nucleoside hydrolase domain (AUL71835.1). Aside from these RNA polymerase subunits, both TV and SMBV release proteins that facilitate transcription. SMBV releases a poly (A) polymerase (AHJ40056.1), an mRNA-capping enzyme (AHJ40083.1), and an anaerobic transcription regulator (AMK61903.1). TV releases an SNF2 family helicase (AUL77941.1), an ATP-dependent RNA helicase (AUL77829.1), and a mimivirus-like elongation factor (AUL78714.1).

Many of the proteins we identified matched proteins that one would expect to be released during the initial stages of viral infection and greatly support our hypothesis that the *in vitro* stages generated in this study are reflective of those that occur *in vivo*. These data provide new insights into GV biology and ultimately lead to our proposed model (see next sections).

SMBV and TV Also Release Novel Proteins during Stargate Opening

SMBV and TV also release proteins that are relatively uncommon among viruses. These proteins include metal-binding homeostasis proteins as well as chemotaxis-regulating proteins. Functions for all predicted proteins are based on sequence similarity to known proteins; we did not confirm the biological function of these proteins with experimental data, and this will require follow-up studies.

Our mass spectrometry data conclusively show that both SMBV and TV release proteins predicted to play a role in maintaining homeostasis (Figure 5E). Many of these proteins likely have redox activity, protecting the virus and its cargo from reactive oxygen species (ROS) that can be found in the host phagosome (Flannagan et al., 2015). These proteins include several thioredoxin-like or thioredoxin-domain-containing proteins (SMBV: AHJ40071.1 and AHJ40129.2; TV: AUL77963.1) and glutaredoxins (SMBV: AMK618100.1; TV: AUL78724.1). TV releases a catalase protein (AUL78097.1) as well as glyoxylase (AUL78134.1) although SMBV releases a prolyl 4-hydroxylase (AMK61959.1). These proteins are also projected to protect the GV from ROS during the infection process. Here, we show that these proteins are indeed released very early in the infection process.

Redox-active proteins are also thought to play an important role in protecting the viruses from the harsh conditions present in the host phagosome. During phagocytosis, amoebal phagosomes drop to \sim pH 4 (not low enough to trigger stargate opening *in vitro*), but they are also inundated with metals (like Cu and Zn) and ROS (German et al., 2013; Lopez and Skaar, 2018). Both viruses release metal-binding proteins (e in Table 1), including SMBV's lanosterol demethylase (AHJ40393.1)—a cytochrome p450-like protein—and prolyl 4-hydroxylase (AMK61959.1) and TV's mg709 (AUL77661.1)—a putative prolyl 4-hydroxylase with iron ion binding capabilities—and Cu-Zn superoxide dismutase (AUL78503.1). In conjunction with the ROS-mitigating proteins described above, these proteins likely allow GVs to survive the onslaught of low pH, high ROS, and high metal concentration found inside of the host phagosomes. We also note that the low pH of the phagosomes is similar to the low pH used in our *in vitro* assay, likely reflecting a physiologically relevant stage that describes GV infection mechanisms.

Although *Tupanvirus* infection is hypothesized to occur through phagocytosis (Abrahão et al., 2018; Silva et al., 2019), no biological data have yet been provided to substantiate said hypothesis. This proposal stems from visualization of phagocytosis of TV by *Vermamoeba vermiformis* and subsequent TV stargate opening via thin-section TEM (Silva et al., 2019). Thin-section TEM, embedding biological samples within epoxy resin then slicing thin, electron translucent sections off of the block, is prone to structural artifacts (Baker et al., 1999). Therefore, it is critical that any hypotheses generated from thin-section TEM imaging are supported by data from another technique. The release of proteins capable of mitigating the harsh environment of the amoebal phagosome provides biological evidence to support this hypothesis.

SMBV and TV also contain proteins that are predicted to regulate chemotaxis. SMBV releases a chemotaxis protein (AHJ40337.1) that shares homology with the putative chemotaxis protein CheD found in mimivirus (AKI80461.1) and TV (AUL78687.1). CheD proteins regulate chemotaxis via deamidation of chemotaxis receptors (InterPro). TV has been shown to shut down host chemotaxis (Oliveira et al., 2019; Rodrigues et al., 2019a), and it is likely that these CheD-like chemotaxis regulation proteins are involved in this process. Although TV does contain a CheD-like chemotaxis protein that was identified in the total virion MS data, this protein was not released following low pH treatment.

Opening the Stargate to New Avenues of GV Research

We were able to mimic four unique stages of the GV genome release process (Figure 6). GV particles that mimic these genome release stages have been seen in previous experiments (Abrahão et al., 2018; Schrad et al., 2017; Xiao et al., 2009; Zauberman et al., 2008) although relied on finding the “one-in-a-million” particle in the correct state. We are now able to mimic GV genome release stages reliably and with high frequency. Additionally, these conditions forgo the need to synchronize infection and trap GV particles in phagosomes at very specific times. Eschewing the host cell may limit specific avenues of study, such as searching for a host receptor(s), but it dramatically simplifies any studies aimed at the

virus and the changes it undergoes during the genome release process.

Additionally, we have identified proteins that are released during the initial stages of infection in two GVs, SMBV and TV. Over half of the proteins released by these viruses are annotated as hypothetical, low complexity, or as an ORFan (which are new protein-coding genes restricted to taxonomically related genomes but are without homologs in other lineages). We were able to provide functional predictions for some of these proteins through homology. The release of these proteins at the initiation of stargate opening suggests that these proteins play an important role in the early stages of GV infection (phagosome survival, genome translocation, early transcription, host defense suppression, etc.). The exact functions of these proteins, as well as how their interactions mediate and orchestrate GV infection, are prime candidates for future study, and the importance is enhanced by the fact that many GVs appear to share similar strategies for genome release. All four of the GVs tested in this study responded similarly to treatment *in vitro*, suggesting that these GVs utilize similar molecular forces during genome release and likely similar proteins.

STAR★METHODS

Detailed methods are provided in the online version of this paper and include the following:

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 - Data and Code Availability
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 - Giant Viruses
- **METHOD DETAILS**
 - Treatment of SMBV Particles and Image Analysis
 - Cryo-Electron Microscopy (Cryo-EM) and Cryo-Electron Tomography (Cryo-ET)
 - Scanning Electron Microscopy
 - Differential Mass Spectrometry
- **QUANTIFICATION AND STATISTICAL ANALYSIS**
 - Mass Spectrometry Analysis

SUPPLEMENTAL INFORMATION

Supplemental Information can be found online at <https://doi.org/10.1016/j.cell.2020.04.032>.

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AUTHOR CONTRIBUTIONS

Designed and Carried out Experiments, J.R.S. and K.N.P.; Wrote the Manuscript, J.R.S.; Data Analysis, J.R.S., J.R.C., and K.N.P.; Viral Sample Preparation, J.R.C.; Manuscript Review, J.R.C., J.S.A., and K.N.P.; Ultrathin Sectioning Imaging, J.S.A.

DECLARATION OF INTERESTS

The authors declare no competing interests.

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STAR★METHODS

KEY RESOURCES TABLE

REAGENT or RESOURCE	SOURCE	IDENTIFIER
Bacterial and Virus Strains		
Samba Virus	Campos et al., 2014	N/A
Tupanvirus soda lake	Abrahão et al., 2018	N/A
Antarctica virus	Andrade et al., 2018	N/A
Mimivirus strain M4	Boyer et al., 2011	N/A
Biological Samples		
<i>Acanthamoeba castellanii</i>	ATCC	ATCC 30010
Chemicals, Peptides, and Recombinant Proteins		
Sodium phosphate, dibasic	Sigma-Aldrich	Cat # S9390
Urea	VWR Life Sciences	Cat # 0568
Guanidinium Hydrochloride	Amresco	Cat # 0118
Triton X-100	Alfa Aesar	Cat # A16046
Chloroform	EMD	Cat # CX1055
Dimethyl sulfoxide	JT Baker	Cat # 9924-01
Bromelain (from pineapple)	Sigma	Cat # 647-005-00-X
Proteinase K	Roche	Cat # 03115879001
DNase I	Roche	Cat # 10104159001
Lysozyme	Amresco	Cat # 0663-5G
712 PYG	ATCC	N/A
Sucrose	JT Baker	Cat # 4072-05
Software and Algorithms		
IMOD	Mastrorade and Held, 2017	https://bio3d.colorado.edu/imod/download.html
SerialEM	Mastrorade, 2005	https://bio3d.colorado.edu/SerialEM/download.html
Leginon	Suloway et al., 2005	http://nramm.nysbc.org/downloads/
Amira	ThermoFischer Scientific	https://www.thermofisher.com/us/en/home/industrial/electron-microscopy/electron-microscopy-instruments-workflow-solutions/3d-visualization-analysis-software/amira-life-sciences-biomedical.html
MaxQuant	Cox and Mann, 2008	https://www.maxquant.org/
Andromeda	Cox et al., 2011	http://coxdocs.org/doku.php?id=maxquant:start/
MotionCor2	Zheng et al., 2017	https://msg.ucsf.edu/software
Other		
Quantifoil R2/2 Grids	Electron Microscopy Sciences	Cat # Q250-CR2
10 nm Nanogold Fiducial Markers	Aldrich (Millipore)	Cat # 741957
VSWP Membrane Filter Discs	Millipore	Cat # VSWP01300
Quantifoil R3.5/1 Grids	Electron Microscopy Sciences	Cat # Q225CR-35
Deposited Data Generated		
pH 2 treated sample, Video S3	This work	EMDB: EMD-20747
100°C treated sample, Video S4	This work	EMDB: EMD-20748
pH 2 + 100°C treated sample, Video S6	This work	EMDB: EMD-20746
pH 2 + 100°C treated sample, Video S8	This work	EMDB: EMD-20745

RESOURCE AVAILABILITY

Lead Contact

Further information and requests for resources and reagents should be directed to and will be fulfilled by the Lead Contact, Kristin Parent (kparent@msu.edu).

Materials Availability

This study did not generate new unique reagents.

Data and Code Availability

The full proteomics datasets generated in this study are available upon request from the Lead Contact.

Three-dimensional tomograms have been deposited to the Electron Microscopy Database (EMDB) under the ID codes EMD-20747 (pH 2, [Video S3](#)), EMD-20748 (100°C, [Video S4](#)), EMD-20746 (pH 2 + 100°C, [Video S6](#)), and EMD-20745 (pH 2 + 100°C, [Video S8](#)).

EXPERIMENTAL MODEL AND SUBJECT DETAILS

Acanthamoeba castellanii

Acanthamoeba castellanii cells were purchased from ATCC (ATCC 30010). *Acanthamoeba castellanii* (ATCC 30010) was cultivated in 712 PYG media w/Additives (ATCC recipe) at pH 6.5 in the presence of gentamicin (15 µg/mL) and penicillin/streptomycin (100 U/mL) at 28°C to reach a 90% confluence.

Giant Viruses

Tupanvirus soda lake (TV), Antarctica virus, and Samba virus (SMBV) were isolated previously ([Abrahão et al., 2018](#); [Andrade et al., 2018](#); [Campos et al., 2014](#)). M4 virus was kindly provided by Dr. Bernard La Scola and Dr. Thomas Klose ([Boyer et al., 2011](#)). *Acanthamoeba castellanii* (ATCC 30010) was cultivated in 712 PYG media w/ Additives (ATCC recipe) at pH 6.5 in the presence of gentamicin (15 µg/mL) and penicillin/streptomycin (100 U/mL) at 28°C to reach a 90% confluence. SMBV or TV virions were diluted in phosphate buffered saline (PBS) and added to the cells to a multiplicity of infection of 5 (TV) or 10 (SMBV). An initial incubation was carried out for one hour at room temperature. After the initial incubation, additional PYG media was added to the cells and the flasks were incubated at 28°C for 48 hours. After 48 hours, more of the free amoebal cells had been lysed. Suspensions containing cell debris and cell particles were centrifuged at 900 x g to pellet residual cells. The resulting supernatant was filtered using a 2 µm filter and was immediately applied to a 22% sucrose cushion (w/w) at 15,000 x g for 30 min. Viral pellets were resuspended in PBS and stored at -80°C. Viruses were tittered using the Reed-Muench protocol ([Ramakrishnan, 2016](#)). On average, virus isolation yielded 10¹⁰ TCID₅₀/mL (TCID = tissue culture infective dose).

METHOD DETAILS

Treatment of SMBV Particles and Image Analysis

Determining the Percentage of Open SMBV Particles

For all treatments, the percentage of open SMBV particles (POP) was determined via 2D cryo-electron microscopy. This was calculated by dividing the number of clearly opened particles by the number of total particles visualized for each condition. In general, these experiments were performed either in duplicate or triplicate in order to generate a minimum of 50 particles imaged per condition. These percentages were compared to the native (untreated) level of spontaneous SMBV particle opening, determined previously to be ~5% ([Schrad et al., 2017](#)).

Conditions That Did Not Increase POP

SMBV particles were treated with various conditions that have been shown to disrupt/destroy other viruses. These conditions include urea, guanidinium hydrochloride, DMSO, Triton X-100, chloroform, DNase I, and an enzyme cocktail (lysozyme, bromelain, proteinase K) that was previously shown to remove APMV fibers ([Kuznetsov et al., 2010](#)). Treatments were applied for 1-2 hours prior to POP determination via cryo-EM. Concentrations for the various conditions, as well as the resultant POP values, can be found in [Table S1](#).

pH Titration of SMBV Particles

25-50 µL of SMBV particles were added to Millipore VSWP Membrane Filter dialysis discs (0.025 µm cutoff) which were then floated onto ~25 mL of 20 mM sodium phosphate buffer, adjusted to the desired pH. The samples were allowed to equilibrate for 1.5-2 hours. For conditions where low pH would interfere with additional treatment (e.g., pH 2 + DNase I or pH 2 samples submitted for mass spectrometry) the particles were dialyzed for an additional 1.5-2 hours against pH 7.0 buffer to restore neutral pH.

High Temperature Incubation

GV particles were incubated in a BioRad T100 thermal cycler at 80, 89, and 100°C for 1 hour. SMBV particles remained intact following 1 hours at 100°C, so additional incubations at 100°C were performed at 2, 3, or 6 hours. As a control, SMBV particles were also incubated for 1 hour at room temperature (25°C).

Combining High Temperature and Low pH

To determine the effect of combining low pH and high temperature, GV particles were sequentially treated with pH 2 and 100°C. First, SMBV particles were dialyzed against 20 mM sodium phosphate buffer, adjusted to pH 2, for 2 hours. Following dialysis, SMBV particles were incubated at 100°C for 3 hours.

Cryo-Electron Microscopy (Cryo-EM) and Cryo-Electron Tomography (Cryo-ET)

Sample Preparation

Samples for cryo-EM and cryo-ET were prepared as described previously (Schrad et al., 2017). Briefly, small (3–5 μL) aliquots of virus particles were applied to R2/2 (cryo-EM) or R3.5/1 (cryo-ET) Quantifoil grids (Electron Microscopy Solutions) that had been plasma cleaned for 20 s in a Fischione model 1020 plasma cleaner. Prior to virus addition, 5–10 μL of 10 nm nanogold fiducial markers were applied to the R3.5/1 grids and were air-dried to provide markers for fiducial alignment of the tilt series. The samples were plunge frozen in liquid ethane using a manual plunge-freezing device (Michigan State University Physics Machine Shop) at room temperature using approximately 5–7 s of blotting time per grid. Frozen-hydrated samples were stored, transferred, and imaged under liquid nitrogen temperatures.

2D Cryo-Electron Microscopy

Cryo-EM experiments were performed at Michigan State University. Virus particles were imaged in either a JEOL 2200-FS TEM or an FEI Talos Arctica. The JEOL was operated operating at 200 keV, using low dose conditions controlled by SerialEM (version 3.5.0-beta, (Mastronarde, 2005)) with the use of an in-column Omega Energy Filter operating at a slit width of 35 eV. Micrographs were recorded at 25 frames per second using a Direct Electron DE-20 direct detector, cooled to -38°C . Motion correction was performed using the Direct Electron software package (Direct Electron, LLC). The Arctica was also operated at 200 keV, under low dose conditions controlled by EPU. Micrographs were recorded on a Falcon 3 direct detector operating in linear mode. Micrographs were collected between 8,000 and 10,000 X nominal magnification (6.87 and 5.30 $\text{\AA}/\text{pixel}$, respectively) on the JEOL and at 17,500 X nominal magnification (6.03 $\text{\AA}/\text{pixel}$) on the Arctica. The objective lens defocus settings ranged from 10 to 15 μm underfocus. Micrographs were collected for 5 s, resulting in a total dose of $\sim 35 \text{ e}/\text{\AA}^2$. For bubblegram imaging, the SMBV particles were imaged for an additional four exposures, resulting in a total does of $\sim 140 \text{ e}/\text{\AA}^2$.

Cryo-Electron Tomography

Cryo-ET tilt series were collected using a Titan Krios TEM operating at 300 keV with a post-column GIF (20 eV slit width) under low dose conditions controlled by SerialEM or Legion at Purdue University. Images were collected using a Gatan K2 direct electron detector operating at 100 ms/frame. Images were collected in super resolution mode between 33,000 and 53,000 X nominal magnification (2.12 – 1.33 $\text{\AA}/\text{pixel}$). Tilt series were carried out between $\pm 50^\circ$ with bidirectional image collection every 2° . Images were collected for 5 s (100 frames/tilt image), resulting in $\sim 2.5 \text{ electrons}/\text{\AA}^2$ per tilt image ($\sim 125 \text{ electrons}/\text{\AA}^2$, total exposure dose).

Individual micrographs were corrected for particle motion and binned by a factor of two using MotionCor2 v1.2.0 (Zheng et al., 2017) and the corrected images were stitched back into a tilt series using the newstack functionality in IMOD (Mastronarde and Held, 2017). Tilt series alignment, using fiducial markers, and tomogram generation was carried out using IMOD v4.7.5. Final tomogram volumes were generated using ten iterations of the SIRT reconstruction method (Mastronarde, 1997) then filtered using the smooth (3x3 kernel) and median (size 3) options in IMOD. Select tomograms were annotated using Amira v2019.2 (ThermoFisher Scientific).

Scanning Electron Microscopy

SEM. Preparation and Imaging

GV particles were imaged using a JEOL JSM-7500F scanning electron microscope. Prior to imaging, virus particles were desiccated using an EM CPD300 critical point dryer, fixed with glutaraldehyde onto poly-L-Lysine treated SEM slides, and sputter coated with a $\sim 2.7\text{nm}$ layer of iridium using a Q150T Turbo Pumped Coater. Particles were imaged between 8,500 X and 85,000 X nominal magnification.

Ultrathin Sections and Transmission Electron Microscopy

A. castellanii cells were infected by APMV or Tupanvirus at a multiplicity of infection of 5 and fixed at 1 hours post infection with 2.5% glutaraldehyde in a 0.1 M sodium phosphate buffer for 1 h at room temperature. The amoebas were postfixed with 2% osmium tetroxide and embedded in Epon resin. Ultrathin sections (50–100 nm) were then analyzed using transmission electron microscopy (Spirit Biotwin FEI-120 kV) at the Center of Microscopy of Universidade Federal de Minas Gerais (UFMG).

Differential Mass Spectrometry

Sample Preparation

SMBV and TV particles were dialyzed against 20 mM sodium phosphate buffer, adjusted to pH 2, for 2 hours, as described above. An aliquot of each virus was left undialyzed as a control (Material Applied, MA). Following dialysis, proteins that had been released from the viral particles were separated from the virions via centrifugation in a microcentrifuge at 8,000 x g for 15 minutes. Visible viral pellets were resuspended in the same volume as the supernatant using 20 mM sodium phosphate buffer, adjusted to pH 7.0. Two technical replicates were created for each sample. An aliquot of each sample was used for SDS-PAGE.

Each sample was TCA precipitated and submitted for LC/MS/MS analysis to the MSU Proteomics Core. Prior to submission, samples were run on a 15% polyacrylamide gel at a voltage of 200 V for 45 minutes. TV and SMBV gel bands visible by Coomassie blue stain were excised and submitted for MS analysis as well.

Proteolytic Digestion

TCA precipitated pellets were re-suspended in 270 μ L of 100mM ammonium bicarbonate supplemented with 10% trifluoroethanol. Samples were reduced and alkylated by adding TCEP and Iodoacetamide at 10mM and 40mM, respectively and incubating for 5min at 45C with shaking at 1400 rpm in an Eppendorf ThermoMixer. Trypsin, in 100mM ammonium bicarbonate, was added at a 1:100 ratio (wt/wt) and the mixture was incubated at 37C overnight. Final volume of each digest was \sim 300 μ L. After digestion, the samples were acidified to 2% TFA and subjected to C18 solid phase clean up using StageTips (Rappsilber et al., 2007) to remove salts.

LC/MS/MS and Data Analysis

An injection of 5 μ L was automatically made using a Thermo EASYnLC 1200 onto a Thermo Acclaim PepMap RSLC 0.075mm x 20mm C18 trapping column and washed for \sim 5min with buffer A. Bound peptides were then eluted over 95min with a gradient of 8%B to 42%B in 84min, ramping to 100%B at 85min and held at 100%B for the duration of the run (Buffer A = 99.9% Water/0.1% Formic Acid, Buffer B = 80% Acetonitrile/0.1% Formic Acid/19.9% Water) at a constant flow rate of 300nl/min. Column temperature was maintained at a constant temperature of 50°C using an integrated column oven (PRSO-V2, Sonation GmbH, Biberach, Germany). Eluted peptides were sprayed into a ThermoScientific Q-Exactive HF-X mass spectrometer using a FlexSpray spray ion source. Survey scans were taken in the Orbi trap (60,000 resolution, determined at m/z 200) and the top ten ions in each survey scan are then subjected to automatic higher energy collision induced dissociation (HCD) with fragment spectra acquired at 7,500 resolution. The resulting MS/MS spectra are converted to peak lists using MaxQuant v1.6.0.1 (Cox and Mann, 2008) and searched using the Andromeda (Cox et al., 2011) algorithm against a protein database containing sequences from SMBV or TV and *Acanthamoeba castellanii* (each downloaded from NCBI, <https://www.ncbi.nlm.nih.gov>). Common laboratory contaminants were included in the Andromeda search. Protein and peptide FDR for all searches were set to 1%.

Mass Spectrometry Data Synthesis

The percentage of the total LFQ signal each protein was responsible for in each sample was calculated by dividing the individual protein LFQ signal by the total LFQ signal for the sample, excluding contaminants. Proteins that are released from the viral particles are expected to make up a higher percentage of the supernatant sample than the whole virion (MA), so the ratios of these two percentages were calculated (Tables 1 and S3). Proteins with a supernatant:MA ratio > 1 were selected for further analysis.

Classification/Functional Annotation of Proteins Identified via MS

TV and SMBV proteins released at low pH were classified via their predicted functions and domains. Primary functional annotation had been carried out previously for both TV (Abrahão et al., 2018) and SMBV (Campos et al., 2014). Additional functional prediction, as well as homology prediction between the two viruses, was carried out through the use of the NCBI BLAST database (NCBI) as well as the HHBLITS server (Remmert et al., 2011) and the InterPro database (Mitchell et al., 2019). Domain prediction was carried out by searching the InterPro database and utilizing the PSIPRED server (Buchan and Jones, 2019) with the DISOPRED3 (Jones and Cozzetto, 2015) functionality activated.

QUANTIFICATION AND STATISTICAL ANALYSIS

Mass Spectrometry Analysis

LFQ intensities for both SMBV and TV spectra were detected in triplicate. For each virus, the initial run did not produce high quality data so these intensities were disregarded. LFQ intensities from the remaining two runs were averaged together to produce the reported intensity (Table S3).

Supplemental Figures

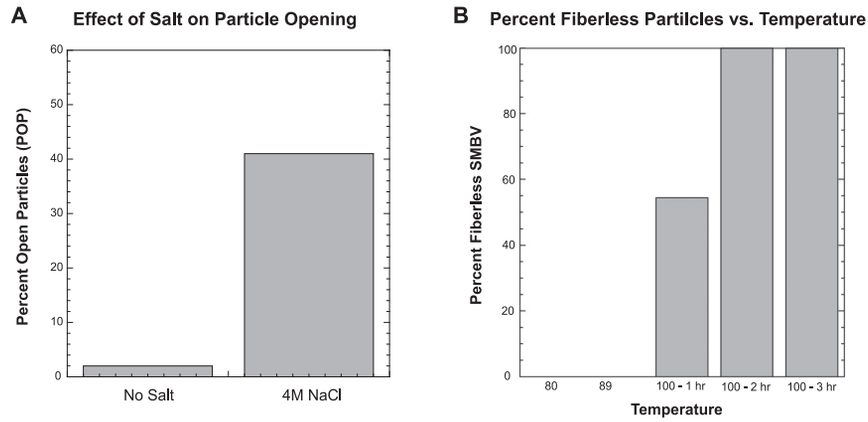


Figure S1. Percentage of Fiberless SMBV Particles at Varying Temperatures, Related to Figure 1

A) Histogram of the percentage of open SMBV particles after treatment with salt. B) Histogram of the percentage of fiberless (open or unopen) SMBV particles at various temperatures and incubation times.

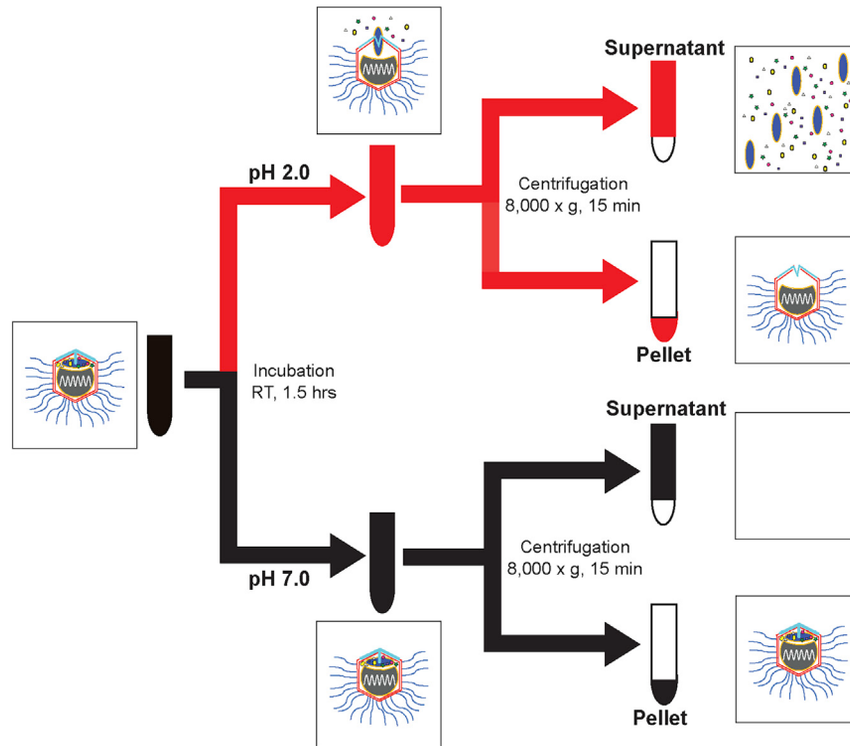


Figure S2. Sample Preparation for SDS-PAGE and LC/MS/MS Experiments, Related to Figures 4 and 5
A cartoon of the workflow schematic used to prepare samples for both the SDS-PAGE and LC/MS/MS experiments.