

1 **Cryo-electron tomography reveals coupled flavivirus replication, budding and maturation**

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34 **Abstract**

35 Flaviviruses replicate their genomes in replication organelles (ROs) formed as bud-like invaginations
36 on the endoplasmic reticulum (ER) membrane, which also functions as the site for virion assembly.
37 While this localization is well established, it is not known to what extent viral membrane remodeling,
38 genome replication, virion assembly, and maturation are coordinated. Here, we imaged tick-borne
39 flavivirus replication in human cells using cryo-electron tomography. We find that the RO membrane
40 bud is shaped by a combination of a curvature-establishing coat and the pressure from intraluminal
41 template RNA. A protein complex at the RO base extends to an adjacent membrane, where immature
42 virions bud. Naturally occurring furin site variants determine whether virions mature in the immediate
43 vicinity of ROs. We further visualize replication in mouse brain tissue by cryo-electron tomography.
44 Taken together, these findings reveal a close spatial coupling of flavivirus genome replication,
45 budding, and maturation.

46

47 **Introduction**

48 *Orthoflaviviruses* (henceforth flaviviruses) are a large genus of arthropod-borne, positive-sense RNA
49 viruses within the *Flaviviridae* family. The mosquito-borne Dengue virus alone is estimated to yearly
50 cause hundreds of millions of human infections, some progressing to the severe condition known as
51 dengue shock syndrome¹. Human infections with tick-borne flaviviruses are less frequent, but can
52 have severe outcomes. Tick-borne encephalitis virus (TBEV) is the namesake virus of the “TBEV
53 serocomplex” which includes other tick-borne flaviviruses such as Powassan virus and the low-
54 pathogenic Langat virus (LGTv). Pathogenic tick-borne flaviviruses have a strong neurotropism in
55 mammals, and can cause encephalitis with debilitating or deadly outcome in humans².

56 After entering the cell through endocytosis, the flavivirus genome is translated as a single,
57 transmembrane polyprotein, which is subsequently cleaved by host and viral proteases into ten
58 individual proteins. Seven of these are the non-structural (NS) proteins, which serve to replicate the
59 viral genome. Of the NS proteins, NS3 and NS5 are cytoplasmic enzymes that serve as protease and
60 helicase (NS3), and RNA-dependent RNA polymerase and methyl transferase (NS5). The remaining
61 NS proteins include the endoplasmic reticulum (ER) lumen-resident peripheral membrane protein
62 NS1, and the integral membrane proteins NS2A, NS2B, NS4A and NS4B. Viral genome replication
63 takes place on a transformed, dilated ER containing multiple bud-like membrane invaginations^{3,4}.
64 These invaginations, referred to as replication organelles (ROs), are the site of viral RNA replication⁵⁻
65 ⁹. RO-like membrane rearrangements can be formed by a subset of NS proteins even in the absence of
66 viral RNA replication^{5,10,11}, but require interactions with host ER proteins^{5-9,12,13}. Electron microscopy
67 of resin-embedded, infected cells has shown that the RO is a 80-90 nm, near-spherical bud with a ~10
68 nm opening towards the cytoplasm¹⁴⁻¹⁷. However, due to the destruction of protein structure by resin
69 embedding, the organization of proteins and RNA in the RO is still unknown. Virion assembly also
70 takes place at the ER, when a cytoplasmic complex of viral RNA and C protein interacts with the
71 transmembrane envelope proteins prM and E, followed by budding into the ER lumen. NS2A has been
72 suggested as a key viral protein coupling replication and assembly¹⁸⁻²⁰, and resin-embedding electron
73 microscopy has visualized putative virions in the immediate vicinity of ROs^{14,16,21}. Newly formed,
74 immature virions have a spiky surface covered with extended prM-E trimers²²⁻²⁴. The cleavage of prM
75 by the host-cell protease furin, which is thought to occur in the trans-Golgi, leads to a structurally
76 rearranged, infectious, mature virion with smoother appearance^{25,26}. If virion assembly and maturation
77 are directly linked to ROs is currently unknown.

78 To shed light on the interactions between flavivirus replication, assembly and maturation, we
79 performed *in situ* cryo-electron tomography²⁷⁻³¹ on human cells and mouse brain tissue infected with
80 LGTV, and a novel, chimeric LGTV carrying TBEV structural proteins. The data suggest a mechanism
81 for RO membrane remodeling, the presence of a protein complex tethering the RO membrane to an
82 apposed ER membrane, and a close proximity of virion assembly and maturation.

83

84

85 **Results**

86 **Cryo-electron tomography reveals two states of replication organelles in Langat virus-infected
87 cells**

88 To explore the macromolecular architecture of flavivirus ROs, we grew human A549 cells on EM
89 grids and infected them with LGTV. Cells were plunge-frozen at 24 h post infection (h.p.i) and
90 subjected to focused-ion-beam milling, after which lamellas containing the infected cytoplasm were
91 imaged using cryo-electron tomography (cryo-ET) (Table 1). The tomograms revealed a dilated ER
92 inclusive of clustered ROs (Fig. 1 and Supplementary video 1). ROs were clearly identified as near-
93 spherical membrane invaginations into the ER lumen, of a kind not present in uninfected cells (Fig.
94 S1A-B). The vicinity of the remodeled ER contained *bona fide* ribosomes as well as mitochondria
95 immediately apposed to the ER membrane (Fig. 1A-D). ROs frequently appeared in clusters within the
96 lumen of dilated ER, as in Fig. 1A-D in which a single, dilated ER cisterna contained >10 ROs within
97 the field of view (bearing in mind that the RO cluster probably extended beyond the depth of the
98 lamella). The same ER cisterna additionally contained a virus particle with the characteristic spiky
99 appearance of immature flaviviruses (Fig. 1A-B, orange arrow, and Fig. 1D). The majority of ROs
100 contained filamentous densities, presumably the replicating double-stranded form of the viral RNA,
101 within their lumen (Fig. 1A-D). On the other hand, several ROs were devoid of internal filamentous
102 structures (Fig. 1A-C, white arrows). These two types of ROs will henceforth be denoted as ‘filled’
103 and ‘empty’, respectively (Fig. 1E). In 23 tomograms, $83\pm14\%$ of ROs were filled (Fig. 1F). The
104 empty ROs were significantly smaller with an average diameter of 46 ± 8 nm ($N=25$), compared to the
105 filled ROs at 85 ± 5 nm ($N=63$) (Fig. 1G). In summary, we established a workflow to image flavivirus
106 replication by cryo-ET, revealing that ROs exist in two forms: with and without luminal filamentous
107 densities.

108

109 **A combination of membrane coat and RNA-induced pressure determines RO morphology**

110 ROs of other viruses depend on the pressure from intraluminal double-stranded RNA (dsRNA) to
111 inflate and stabilize the curved RO membrane³². The presence of empty LGTV ROs speaks against
112 this mechanism for flaviviruses, and we thus reasoned that a coat might confer a spontaneous
113 curvature to the RO membrane. To investigate whether a curvature-inducing coat is present on ROs,
114 we assessed if RO membrane thickness is in line with a layer of surface-bound protein layer. Indeed,
115 by visual inspection of tomograms, the membranes of both empty and filled ROs appeared thicker than
116 the surrounding ER membrane (Fig. 2A). No distinct, repeating macromolecules were visible on the
117 RO membranes. While this does not exclude the presence of a protein coat composed of smaller or
118 membrane-integral proteins, it does make its characterization by subtomogram averaging less likely to
119 be successful. Instead, we extended our previously developed surface morphometrics toolbox to allow
120 for local estimation of membrane thickness³³. This software allows calculation of the average
121 membrane thickness within a user-selected area, based on average density profiles normal to the
122 membrane. Both for single ROs and their surrounding ER membrane, this yielded reliably
123 interpretable density profiles (Fig. 2B-C). Color coding membranes by thickness indicated that RO
124 membranes are consistently thicker than the surrounding ER membrane (Fig. 2D). Indeed, in four
125 tomograms we measured a significant difference in membrane thickness with the ER membrane being
126 3.4 ± 0.2 nm ($N=4$), and RO membranes 4.0 ± 0.2 nm ($N=132$) (Fig. 2E). On the other hand, the RO
127 membrane thickness appeared largely independent of RO size, as estimated from their radii of
128 curvature (Fig. 2F).

129 The observation that flavivirus ROs can form without detectable luminal dsRNA, and the consistent
130 presence of a membrane coat, distinguish them from alphavirus ROs that have a near-identical
131 membrane shape. Based on cryo-ET data, we recently published a mathematical model of alphavirus

132 RO membrane budding, which showed that the pressure from intraluminal dsRNA, together with
133 constraint of the membrane neck, is sufficient for the creation of the RO membrane bud³². We next
134 adapted this mathematical model to explain flavivirus RO membrane remodeling. We assume that the
135 membrane coat generates a spontaneous curvature H_0 of the RO membrane. Such a spontaneous
136 curvature can be generated by the protein structure as well as by crowding³⁴. Furthermore, we consider
137 that in ROs that enclose dsRNA, the RNA exerts a pressure P on the membrane. The total energy E of
138 the RO membrane is then composed of an integral over the membrane surface A and the contribution
139 of the pressure, which scales with the volume V ,

140
$$E = \int_A dA [2\kappa(H - H_0)^2 + \sigma] - PV,$$

141 (1)

142 where the first term describes the bending energy according to the Helfrich model, with κ the bending
143 stiffness and H the mean curvature³⁵. The second term in Eq. 1 contains the membrane tension σ .
144 Motivated by the experimentally observed shapes, we describe the ROs as spheres with a radius R , as
145 schematically depicted in Fig. 2G, simplifying Eq. (1) to

146
$$E = 8\pi\kappa(1 - H_0R)^2 + 4\pi\sigma R^2 - \frac{4}{3}\pi PR^3.$$

147 (2)

148 However, there are two unknown factors in Equation (2), the spontaneous curvature H_0 and the
149 pressure P in the RO. Based on our observations above, the membrane coat can be assumed to be
150 comparable for empty and filled ROs. Thus, we can take advantage of the imaging of empty ROs to
151 obtain H_0 at vanishing pressure, $P = 0$. Minimizing Eq. (2) with respect to R , we obtain

152
$$H_0 = \frac{1 + \sqrt{1 - 2\frac{\sigma R^2}{\kappa}}}{2R}$$

153 (3)

154 From the experiments we have obtained an average diameter of the empty ROs to be $2R = 46\text{ nm}$. By
155 using previously estimated parameters³² for the membrane properties, i.e., $\sigma = 10^{-5}\text{ N/m}$ and $\kappa =$
156 $10k_B T$, we predict the spontaneous curvature to be $H_0 = 0.04\text{ nm}^{-1}$, which corresponds to a radius of
157 curvature $1/H_0 = 25\text{ nm}$. Next, we want to predict the influence of the pressure generated by the RNA
158 on the spherule size. Since the RNA does not affect the spontaneous curvature H_0 , we keep the above
159 prediction and minimize Eq. 2 with respect to R , giving

160
$$P = \frac{2(2\kappa H_0^2 R - 2\kappa H_0 + \sigma R)}{R^2}$$

161 (4)

162 Now including the predicted $H_0 = 0.04\text{ nm}^{-1}$ with the measured average diameter of RNA-filled ROs
163 $2R = 85\text{ nm}$, we obtain the pressure $P = 5 \cdot 10^{-4} k_B T \text{ nm}^{-3}$. To interpret this value, we compare it
164 with our previous study of alphavirus ROs, which shows that a dsRNA with a length of 2,000-10,000
165 base pairs generates an internal pressure of $10^{-4} - 10^{-3} k_B T \text{ nm}^{-3}$ (see Materials and Methods,
166 section Estimating RO intraluminal pressure).

167 Taken together, flavivirus ROs have a consistently thicker membrane than the surrounding ER,
168 consistent with the presence of small, curvature-stabilizing proteins that set a baseline RO size in the

169 absence of luminal RNA. The size increase from empty to filled ROs is consistent with a single copy
170 of the genome in dsRNA form.

171

172 **Virions form and undergo maturation in the immediate vicinity of replication organelles**

173 In our cryo-electron tomograms we consistently observed virions in the vicinity of ROs, underscoring
174 the strong association between replication and virion assembly (Fig. 3A-C). We next wished to use the
175 structure preservation in cryo-ET to study the spatial relation between virion budding and maturation.
176 In the tomograms, virus particles at different stages of maturation were distinguishable: spiky particles
177 corresponding to immature virions as well as smooth, mature virions (Fig. 3A-C). Subtomogram
178 averaging on a small number of virions confirmed the distinct morphology of the immature and
179 mature virions (Fig. 3D-E), and a good match between the *in situ* averages and structures of purified
180 flaviviruses (Fig. S3). The tomograms also included examples of what seemed to be nearly or recently
181 completed virion budding (Fig. 3A-B). In such events, immature virus particles could be observed
182 right at the membrane, across from ROs (Fig. 3A, orange arrow). Both immature and mature virions
183 were consistently found close to ROs, in separate but intertwined membrane compartments (Fig. 3A-
184 C). While immature and mature particles were not observed in the same membrane compartment,
185 seemingly discrete compartments may have been connected beyond the limited thickness of the
186 lamella. To quantitate the relation of immature and mature particles to ROs, we measured the center-
187 center distance of virions to ROs in 5 tomograms. Immature virions were 95 ± 56 nm (N=37), and
188 mature virions 147 ± 41 nm (N=24) from the closest RO (Fig. 3F). The small but significant distance
189 difference ($p=0.0003$, unpaired t test), together with the observation that immature and mature virions
190 are present in separate compartments, indicates that virion maturation is coupled to a slight spatial
191 separation from ROs but does not necessitate longer-range trafficking. To further investigate the
192 spatial relationship between genome replication, virion assembly and the Golgi apparatus, we
193 performed immunofluorescence light microscopy on fixed, LGTV-infected cells. Furin exhibited co-
194 localization with the Golgi marker GM130 in both uninfected (Fig. S4A-D) and LGTV-infected cells
195 (Fig. 3G-J). Meanwhile, the dsRNA signal was detected in close proximity to and partially
196 overlapping with clusters enriched in furin and GM130 (Fig. 3J). These results corroborate the cryo-
197 ET, reinforcing the close proximity between viral RNA replication, LGTV assembly and maturation
198 within the infected cell. In summary, both virion assembly and maturation can occur in the immediate
199 proximity of ROs, in distinct but intertwined compartments.

200

201 **The TBEV furin site variants R86 and Q86 differ in replication organelle-proximal virion 202 maturation**

203 In purified virions, the transition from immature to smooth conformation can be brought about by pH
204 change even in the absence of furin cleavage. Thus, we wanted to test if the observed RO-proximal
205 virion maturation was dependent on the furin site in prM. To do so, we took advantage of our recently
206 characterized chimeric LGTV, which carries the structural proteins prM and E(ectodomain) from
207 TBEV (Fig. 4A). This virus, rLGT^{T:prME}, is genetically stable and has a low pathogenicity similar to
208 that of wildtype LGTV³⁶. The rLGT^{T:prME} prM and E come from the human TBEV isolate 93/783 of
209 the European subtype, whose prM has an unusual arginine (R) in position 86 where most other strains
210 have a glutamine (Q) (Fig. 4A). This residue is located at the P8 position of the furin cleavage site in
211 prM, and previous studies of flaviviruses has shown that modifications here might influence prM
212 cleavage and virus export³⁷. We thus reasoned that R86 may affect furin cleavage efficiency, and
213 provide a naturally occurring tool to study maturation. Hence, we also produced a version of
214 rLGT^{T:prME} with the more common glutamine 86. These chimeric viruses, henceforth referred to as
215 R86 and Q86, only differ in this amino acid residue. Both viruses replicated with similar kinetics in

216 A549 cells (Fig. S5A), but the R86 virus had a slightly higher lethality in immunocompromised *Ips1^{-/-}*
217 (IFN-β promoter stimulator 1) mice (5 of 5 mice died with R86, 7 of 10 died with Q86, Fig. 4B). No
218 difference in neurovirulence was detected (Fig. 4C). We next wanted to investigate whether the
219 pathogenicity of R86 as compared to Q86 correlated with different prM cleavage kinetics. In a
220 cleavage assay with a peptide corresponding to residues 81-94 of prM, the R86 sequence was cleaved
221 faster by furin than Q86 (Fig. 4D). We noted that the R86 sequence generates a putative second,
222 minimal recognition site (KR) for other proprotein convertases such as PC1/3 and PC2³⁸. In a peptide
223 cleavage assay, the R86 sequence was also cleaved faster than Q86 by PC1/3. However, the cleavage
224 was still completely dependent on the furin recognition site (RTRR), i.e. the putative second PC1/3
225 cleavage site K85-R86 was not sufficient for cleavage by PC1/3 (Fig. 4D). Next, we looked for the
226 presence of unprocessed prM protein in cell supernatant, which would indicate release of immature
227 virus particles. For both viruses, the bulk of cell-bound M was in the form of uncleaved prM, whereas
228 most M in the supernatant was cleaved (Fig. 4E). Interestingly, the R86 chimeric virus had a
229 significantly lower percentage of uncleaved prM in supernatant at 48 h post infection (Fig. 4F),
230 suggesting that R86 confers a more efficient particle maturation. The biochemical and cell assays thus
231 converge on the interpretation that the R86 sequence variant confers faster furin cleavage.

232 Having characterized the different rates of furin cleavage, we then returned to the question of
233 individual virion conformation inside infected cells. We infected cells with R86 and Q86 chimeric
234 viruses and recorded cryo-electron tomograms of the infected cytoplasm as for wildtype LGTV. Both
235 for R86 and Q86, the tomograms showed a similar overall appearance as for wildtype LGTV,
236 including an abundance of filled and empty replication organelles as well as new virions inside dilated
237 ER compartments (Fig. 4G-H, Fig. S5B). In the R86 tomograms, both immature and mature virions
238 were seen, whereas immature virions appeared to predominate in the Q86 tomograms. We calculated
239 the percentage mature virions in a set of tomograms for wildtype LGTV and the chimeric viruses.
240 Whereas the number of virions in individual tomograms was sometimes small, the average fractions
241 mature particles over several tomograms showed a clear trend (Fig. 4I). For Q86, 2.5±5.9% of virions
242 were mature (N=7 tomograms), whereas 46±46% of R86 virions were mature (N=7 tomograms), a
243 significantly higher percentage (p=0.02, unpaired t test). Wildtype LGTV was intermediate to the two
244 recombinant viruses, with 14±25% mature virions. Taken together, a single residue in the distal part of
245 the TBEV furin site affects prM cleavage rates by furin and PC1/3, mean survival in
246 immunocompromised mice, and particle maturation in areas near ROs.

247

248 **A protein complex connects the replication organelle to an apposed ER membrane**

249 A recurring feature in the cryo-electron tomograms was the close proximity of a second ER membrane
250 to the ER membrane containing the ROs (Figs. 1-2). We noticed that this was consistently mediated by
251 a protein complex present at the membrane neck of the replication organelle, connecting this
252 membrane to the adjacent, second ER membrane (Fig. 5A-D). While limited occurrences of these
253 complexes hindered structural analysis, they consistently appeared at the necks of both filled and
254 empty ROs (Fig. 5A-D). From volume estimates of individual complexes, we estimate their molecular
255 masses to 500±151 kDa (Fig. 5E). Interestingly, we observed similar-looking complexes connecting
256 the replication organelle to the site of immature particle budding in the neighboring ER cisterna (Fig.
257 5F-H). This observation suggests that this protein complex might play a role in coordinating the
258 packaging of newly synthesized viral RNA into nascent immature virions.

259

260 **Cryo-ET reveals structural signatures of LGTV replication in *ex vivo* mouse brain**

261 We next wished to explore if the structural features of LGTV replication that we observed in cell lines
262 can also be identified directly in a complex, infected tissue. To do so, we proceeded to set up a

263 workflow for cryo-ET of LGTV-infected mouse brain tissue. We based our approach on our recent
264 publication, in which we imaged entire, *ex vivo*, LGTV-infected brains from type I interferon receptor
265 knockout (*Ifnar*^{-/-}) mice using fluorescence optical projection tomography³⁹. In the 3D volumes of
266 infected brains, immunofluorescence staining against LGTV NS5 protein was particularly strong in the
267 choroid plexus (ChP) regions of the brain (Fig. 6A). ChP is an anatomical substructure responsible for
268 secreting cerebrospinal fluid (CSF) into the ventricles, and as such it interfaces both with the blood
269 and the CSF-filled ventricles (Fig. 6A). The ventricular side of the CSF-producing ependymal cells is
270 covered with cilia and microvilli (Fig. 6A). Based on the consistently strong infection of the ChP, we
271 developed a cryo-ET workflow for imaging ChP that was surgically removed from infected brains *post*
272 *mortem*. Due to the thickness of this sample, we opted for vitrification through high-pressure freezing.
273 The vitrified tissue was trimmed using a cryo-ultramicrotome after which lamellas were milled in
274 place and subjected to cryo-ET (Fig. 6A). The tomograms of ChP revealed a multitude of subcellular
275 structures such as mitochondria, nuclear pore complexes and a centriole (Fig. S6A-B). In addition,
276 some tomograms contained virus-related features consistent with the observations in infected cell
277 lines. This included mature virus particles encapsulated in vesicles close to membranes containing
278 several ROs (Fig. 6B-D). Another tomogram contained a *bona fide* empty RO with seemingly thicker
279 membrane than its limiting ER membrane, mirroring the features seen in cellular tomograms (Fig. 6E-
280 F). Taken together, we present the first cryo-ET data on virus replication in brain tissue. The data
281 support our cell line-based findings of RO-proximal virion maturation, and the presence of empty
282 ROs, and provide a proof of principle that neurotropic virus replication can be studied by cryo-ET
283 directly in infected brain samples.

284

285 Discussion

286 Here, we present an *in situ* structural study of tick-borne flavivirus replication, using cryo-ET of
287 infected cells and mouse brains. Flaviviruses are part of the vast phylum *Kirrinoviricota*, which is
288 characterized by ROs housed in membrane buds⁴⁰. These viruses thus need to encode mechanisms for
289 remodeling host-cell membranes into a high-curvature bud, which is a high-energy and normally
290 transient membrane shape. However, the viruses need to stabilize this bud-shaped membrane
291 throughout hours of viral RNA replication. We recently showed that another genus of *Kirrinoviricota*,
292 alphaviruses, stabilize their RO membrane through a coat-free mechanism that involves bud neck
293 constraint by a viral protein complex, and inflation of the membrane bud by the intraluminal pressure
294 from the viral RNA³². With this mechanism, the size of the replication organelle is determined by the
295 amount of encapsulated RNA, and there are no membrane buds in the absence of intraluminal RNA.
296 Here, we show that flaviviruses employ a different mechanism to shape the RO membrane. A
297 membrane coat establishes a baseline RO size, allowing for the existence of ROs without intraluminal
298 RNA. The tomograms do not indicate an ordered protein coat lattice on the RO membrane, nor clear
299 individual protein densities, but it is possible that the curvature-generating proteins are too small
300 and/or irregularly distributed to be detected by cryo-ET. We suggest that a strong candidate for this RO
301 curvature generator is the 39 kDa, ER lumen-resident, non-structural protein NS1. NS1 is a peripheral
302 membrane protein that has been reported to remodel liposomes⁴¹, and reshape the ER membrane when
303 overexpressed on its own⁴². The increase in RO size due to intraluminal RNA was back-calculated to
304 stem from the pressure of a single viral genomic RNA copy in dsRNA form (Fig. 2G, Fig. S2).
305 Alphavirus ROs also contain a single genome copy, indicating that this may be a conserved feature
306 across *Kirrinoviricota*³². Whether the smaller, empty ROs represent assembly intermediates,
307 disassembly intermediates, or dead-end, failed RO assembly events remains to be determined.

308 A close coupling of flavivirus genome replication and particle budding has been suggested by several
309 lines of evidence^{14,18-20}. Our cryo-electron tomograms clearly resolved the maturation state of individual
310 virions, as supported by the good agreement between the cellular subtomogram averages and structures
311 of purified virions^{23,25} (Fig. 3D-E, Fig. S4). The tomograms frequently revealed immature virions in the

312 immediate vicinity of ROs, sometimes in membrane compartments opposite to ROs (Figs. 3,6). The
313 observation of a ~500 kDa protein complex, connecting the membranes from which ROs and virions
314 form (Fig. 6), shows that flavivirus ROs have a “crown” or “neck complex” akin to those identified for
315 e.g. coronaviruses, nodaviruses and alphaviruses^{30,32,43-50}. Indeed, of the seven flavivirus non-structural
316 proteins, four are integral membrane proteins of unknown structure and no known enzymatic function.
317 Thus, it is possible that these proteins serve a structural role in organizing a membrane-connecting neck
318 complex that coordinates replication and assembly, but larger tomographic data sets would be required
319 to get a decisive subtomogram average from infected cells. Recent publications have highlighted the
320 potential in small-molecule antivirals that target non-enzymatic functions of flavivirus NS proteins⁵¹⁻⁵³.
321 These antivirals were discovered without structural insights into their target proteins, but it is possible
322 that the neck complex we identify here is their target. Either way, a more detailed understanding of the
323 neck complex structure and function may aid the design of improved antiviral strategies.

324 Contrary to the prevailing model, we observed that furin-dependent virus maturation takes place in the
325 immediate vicinity of ROs (Fig. 3). Thus, the entire sequence replication-assembly-maturation is more
326 closely colocalized than previously thought. The maturation compartments are distinct but intertwined
327 with ROs (Fig. 3), suggesting a revised model of flavivirus maturation, in which a virus-induced
328 reorganization of the secretory pathway places Golgi-like maturation compartments in the immediate
329 proximity of ROs. Studying naturally occurring furin site variants, we could show that a single residue
330 in the distal cleavage site affects the RO-proximal virion maturation, while having no effect on virus
331 release and only a minor effect on lethality in an immunocompromised mouse model (Fig. 4). This
332 suggests that the replication and infection of TBEV is robust to variations in its maturation pathway. A
333 further step towards bridging structural and organismal studies of flavivirus replication is taken by the
334 workflow we present for cryo-ET of infected *ex vivo* mouse brain tissue. Tomograms of infected choroid
335 plexus revealed clear structural signatures of ROs and clustering mature virions, similar to those in cell
336 lines (Fig. 7). While cryo-ET has recently been used to study Alzheimer’s disease in human brain⁵⁴, our
337 data are, to the best of our knowledge, the first cryo-ET visualization of infection processes in the brain.
338 Future incorporation of novel lift-out and serial milling techniques into this workflow will allow for
339 faster acquisition of larger cryo-ET data sets on infected brains^{55,56}. This may e.g. enable structural
340 analysis of virion maturation in brain samples, and comparison of replication features between different
341 knock-out mice. In conclusion, our study identifies several novel structural features of tick-borne
342 flavivirus replication, and places them in a cellular context that reveals a high degree of spatial
343 coordination of genome replication, virion assembly and virion maturation.

344

345 Materials and Methods

346 Cell line and culturing

347 The human A549 lung epithelial cell line was grown in DMEM medium supplemented with 10% fetal
348 bovine serum (FBS) and Penicillin Streptomycin GlutaMAX Supplement (Gibco) at 37°C in a 5%
349 CO₂ environment.

350

351 Sample preparation for cryo-electron tomography of cells

352 Ultrafoil Au R2/2 200 mesh grids (200 mesh, Quantifoil Micro Tools GmbH) were glow discharged.
353 Under laminar flow the grids were dipped in ethanol before being placed in μ-Slide 8 Well Chamber
354 (IBIDI) wells. DMEM medium with 10% FBS was added to each well and incubated while cells were
355 being prepared. Fresh medium was added to wells and cells were seeded out at 1.5x10⁴ cells/well. The
356 seeded cells were then placed in a 37°C, 5% CO₂ incubator for 24 h. The medium in the wells was then
357 replaced with serum-free DMEM and either Langat virus wildtype (LGT), recombinant chimeric

358 R86, or recombinant chimeric Q86 added at an MOI of 20. The cells were incubated for one hour
359 before replacing the medium with 2% FBS in DMEM and left to incubate for 24 h. The medium was
360 replaced with fresh DMEM including 2% FBS before being taken for freezing. Plunge freezing into an
361 ethane/propane mix was performed with a Vitrobot (ThermoFisher Scientific) at 22°C, 100%
362 humidity, blotting time of 5 s, and blot force of -5.

363

364 **Preparation of cryo-lamellas of cells**

365 Lamellas were milled from plunge-frozen cells using the Scios dual beam FIB/SEM microscope
366 (ThermoFisher Scientific). Samples were first coated with a platinum layer using the gas injection
367 system (GIS, ThermoFisher Scientific) operated at 26°C and 12 mm working distance for 10 seconds
368 per grid. Lamella were milled at an angle range of 16°-20°. The cells were milled stepwise using a
369 gallium beam at 30kV with decreasing current starting at 0.5 nA for rough milling and ending at 0.03
370 nA for final polishing of the lamella. Lamellas were milled to a nominal 200 nm thickness and stored
371 in liquid nitrogen for less than a week before being loaded into a Titan Krios (ThermoFisher
372 Scientific) for data collection.

373

374 **Cryo-ET data collection on cells**

375 Data were collected using a Titan Krios (ThermoFisher Scientific) at 300 kV in parallel illumination
376 mode. Tilt series acquisition was done using SerialEM⁵⁷ on a K2 Summit detector (Gatan, Pleasanton,
377 CA) in super-resolution mode. The K2 Summit detector was fitted with a BioQuantum energy filter
378 (Gatan, Pleasanton, CA) operated with a 20eV width silt. Areas to be imaged were selected from low-
379 magnification overview images based on the presence of convoluted cytoplasmic membranes. Tilt
380 series were collected using a 100 μm objective aperture and a 70 μm condenser 2 aperture, after
381 coma-free alignment done using Sherpa (ThermoFisher Scientific). Tilt series were collected using the
382 dose-symmetric scheme with a starting angle of -13° to account for lamella pre tilt. The parameters
383 used for acquisition were: 33,000x nominal magnification with a corresponding object pixel size of
384 2.145 Å in super-resolution mode, a tilt range of typically -50° to +50°, defocus between -3 μm and -5
385 μm, tilt increment of 2°, and a total electron dose of 110 e/ Å².

386

387 **Image Processing**

388 Motion correction, tilt series alignment, CTF estimation and correction, and tomogram reconstruction
389 was performed as described previously⁵⁸, using MotionCor2⁵⁹ with Fourier binning of 2, IMOD^{60,61},
390 and CTFFIND4⁶². For visualization and segmentation, tomograms were 3 times binned using IMOD,
391 resulting in a pixel size of 12.87 Å. Tomograms were denoised using cryoCARE⁶³ or IsoNet⁶⁴,
392 occasionally combined with a non-local means filter as implemented in Amira (Thermo Fisher
393 Scientific). Segmentation of tomograms was performed in Amira, with initial membrane tracing and
394 segmentation done using MemBrain V2⁶⁵. Subtomogram averages of mature and immature virus
395 particles were incorporated into segmentation using UCSF Chimera⁶⁶. Amira was used for counting of
396 visually recognizable features (empty and filled ROs, immature and mature particles), measurements
397 of the distances between them, and the volume of the neck complex densities. The neck complex
398 volumes were converted to estimated molecular masses assuming assuming 825 Da/nm³⁶⁷.

399

400 **Subtomogram averaging of virions**

401 From tomograms generated with WARP⁶⁸ at 10Å/px object pixel size, immature and mature particles
402 were manually picked based on their clearly distinguishable appearance. 84 immature and 51 mature
403 particles were extracted from 11 and 3 tomograms, respectively, with a box size of 80*80*80 voxels.
404 Subtomogram averaging was done in Dynamo^{69,70}, following the same procedure for both the
405 immature and mature data sets. Initially, all particles were translationally and rotationally aligned to a
406 single, high-contrast particle from the respective data sets, without symmetrization. These C1 averages
407 were manually rotated and saved in UCSF Chimera⁶⁶ to approximately fit Dynamo's icosahedral
408 convention, after which they were used as a template for a Gold-standard alignment with imposed
409 icosahedral symmetry, using Dynamo's Adaptive bandpass Filtering function. Gold-standard Fourier
410 shell correlation curves estimated the resolution at a cutoff of 0.143 to 33 Å for the immature particles,
411 and 80Å for the mature particles, respectively. The final averages were filtered to this resolution and
412 masked using the spherical alignment mask.

413

414 **Estimating RO intraluminal pressure**

415 In recent work³², we demonstrated the relation between the length of an RNA and the volume of the
416 surrounding spherule. Spherules with a volume of $V = 10^3 - 2 \cdot 10^3 nm^3$ contain $4 \cdot 10^3 - 10^4$ base
417 pairs (Fig. S2A), where the number of base pairs N is well described by

418
$$N = \frac{L_0}{l_{bp}} \left[1 + \frac{\sigma R_N^2}{\kappa} 2 \left(\frac{3}{4\pi} \right)^{4/3} \left(\frac{V}{R_N^3} \right)^{4/3} \right], \quad (S1)$$

419 with $L_0 = 333nm$, $\sigma R_N^2/\kappa = 0.02$, $R_N = 9.6nm$ and the length per base pair $l_{bp} = 0.256nm$.
420 Furthermore, a theoretical model was used to determine the relationship between the scaled pressure
421 PR_N^3/κ and the scaled volume V/R_N^3 (Fig. S2B). Combining both results, we obtain a relation between
422 the pressure P and the number of base pairs N , which shows that an RNA with a length of $2 \cdot 10^3 -$
423 10^4 base pairs corresponds to a pressure of $10^{-3} - 10^{-4} k_B T nm^{-3}$ (Fig. S2C).

424

425 **Immunofluorescence staining**

426 Cells were grown on cover glasses and infected with LGTV as for cryo-ET. At 24 h p.i., cells were
427 fixed with 4% formaldehyde for 20 min at room temperature and then rinsed with PBS. The fixed cells
428 were permeabilized with 0.1% Triton X-100 in PBS for 10 min at room temperature and then rinsed
429 with PBS. The cells were then blocked with 2% BSA in PBS containing 0.05% Tween-20 (PBS-T) for
430 1 hour at RT. The cells were then stained with primary antibodies against furin (goat polyclonal,
431 dilution 1:50, Invitrogen), GM130 (mouse monoclonal clone 35/GM130, dilution 1:300, BD
432 Transduction) or dsRNA (mouse monoclonal clone J2, dilution 1:1000, Scicons, Nordic MUBio;
433 conjugated to allophycocyanin, Abcam) and secondary fluorescent antibodies (donkey anti-goat Alexa
434 Fluor 488, dilution 1:1000, Invitrogen; goat anti-mouse Alexa Fluor 568, dilution 1:1000, Invitrogen)
435 and DAPI diluted in blocking buffer for 1 hour each. Fluorescence images were acquired using a Leica
436 SP8 confocal microscope with a HC PL APO 63x/1.4 oil CS2 objective (Leica). Confocal fluorescence
437 images were analyzed using ImageJ Fiji software⁷¹.

438

439 **Chimeric viruses**

440 A detailed description of chimeric LGTV (rLGT^{T:prME}) generation, rescue, and characterization can
441 be found in our recent manuscript³⁶. In brief, the infectious clone of LGTV, strain TP21 (kind gift of
442 Prof. Andres Merits) was used as genetic background into which the prM and ecto-E from TBEV
443 strain 93/783 (GenBank: MT581212.1) was inserted. Point mutation resulting in the R86Q amino acid
444 substitution in the prM protein were introduced by overlapping PCR with primers; For: 5'
445 GGACGCTGTGGAAACAGGAGGCTCACGGACA 3', Rev: 5'
446 TGTCCGTGAGCCTCTGTTCCCACAGCGTCC 3' (Sigma-Aldrich). RNA was generated from
447 linearized DNA by in vitro transcription and transfected into BHK21 cells using Lipofectamine 2000

448 (Invitrogen). Supernatant from transfected cells was passaged once in A549 mitochondrial antiviral
449 signaling protein (*MAVS*^{-/-}) cells, confirmed by sequencing and used for all downstream experiments
450 without further passaging.

451

452 **RNA isolation and qPCR**

453 RNA was extracted from cell supernatant using Viral RNA kit (Qiagen) according to manufacturer's
454 instructions. The elution volume was kept constant and cDNA was subsequently synthetized from 10
455 µl of eluted RNA using high-capacity cDNA Reverse Transcription kit (Thermo Fisher). LGTV RNA
456 was quantified with qPCRBIo probe mix Hi-ROX (PCR Biosystems) and primers recognizing NS3⁷²
457 on a StepOnePlus real-time PCR system (Applied Biosystems).

458

459 **Western blot**

460 At indicated time points, supernatant was collected and A549 cells infected with rLGTVT:prME R86 or
461 Q86 were lysed in 350 µl of lysis buffer (50 mM Tris-HCl pH7.5 + 150 mM NaCl + 0.1% Triton X-
462 100) complemented with 1x protease inhibitor (cComplete™ ULTRA, Roche, Basel, Switzerland) on
463 ice for 20 min. Following lysis, cellular debris was removed by centrifugation at 14,000 g for 10 min
464 at 4°C. Supernatant or pre-cleared cell lysate was mixed with Laemmli buffer to final concentration 1x
465 and boiled at 95°C for 5 min. Proteins were separated by standard SDS-PAGE and transferred to an
466 Immobilon®-P PVDF Membrane (GE Healthcare, Chicago, IL, USA). Blots were blocked overnight
467 at 4°C in blocking buffer (PBS + 0.05 % Tween 20 + 2% Amersham ECL Prime Blocking Reagent;
468 Cytiva), stained with primary antibodies against NS3⁷³ (chicken polyclonal, diluted 1:1500), tubulin
469 (rabbit polyclonal, diluted 1:4000, Abcam-ab6046) or M⁷⁴ (in-house rabbit polyclonal serum, diluted
470 1:500) overnight at 4°C followed by secondary antibodies (goat anti-chicken Alexa-555, donkey anti-
471 rabbit Alexa-647 (dilution 1:2500, Invitrogen, Waltham, MA, USA) for 1 h at room temperature. Blots
472 were visualized on Amersham™ Imager 680 (GE Healthcare).

473

474 **Enzymatic assays**

475 Synthetic peptides (Biomatik) corresponding to the P13 to P'1 residues of prM from TBEV strain Torö
476 (Dabcyl-GRCGKQEGRSRTRG-E(EDANS)) and 93/783 (Dabcyl-GRCGKREGRSRTRG-
477 E(EDANS)) or corresponding peptides with an impaired furin recognition site (Dabcyl-
478 GRCGKREGSRTRAG-E(EDANS), Dabcyl-GRCGKQEGRTRAG-E(EDANS)) was used as
479 substrate. Cleavage efficiency was assayed using an adapted fluorogenic peptide assay⁷⁵. For furin, 3
480 U furin (Thermo Fisher, Waltham, MA, USA) was mixed with 100 µM of substrate in a total volume
481 of 100 µl reaction buffer (100 mM HEPES pH7.5 + 1 mM CaCl₂ + 0.5% Triton X-100) at 30°C for
482 3h. For proprotein convertase 1/3 (PC1/3), 1 µg recombinant human PC1/3 (R&D Systems,
483 Minneapolis, MN, USA) was mixed with 100 µM of substrate in a total volume of 100 µl reaction
484 buffer (25 mM MES pH 6.0 + 5 mM CaCl₂ + 1% (w/v) Brij-35) at 37°C for 1h. The emission at 490
485 nm measured every 3 min and the average rate calculated by linear regression.

486

487 **Virus infection of mice**

488 All animal experiments were conducted at the Umeå Centre for Comparative Biology (UCCB), under
489 approval from the regional Animal Research Ethics Committee of Northern Norrland and the Swedish
490 Board of Agriculture, ethical permit A9-2018, A41-2019, and conducted as described previously^{39,72}.
491 Briefly, *Mavs*^{-/-} mice in C57BL/6 background (kind gift of Nelson O Gekara, Umeå University) were

492 infected by intraperitoneal injection of 10^4 focus-forming units (FFU) or intracranial injection of 10^2
493 FFU of rLGT^{T:prME} R86 or Q86 diluted in PBS. *Ifnar*^{-/-} mice were intracranially inoculated with 10^4
494 FFU of LGTV and sacrificed when they developed either one pre-defined severe sign or at least three
495 milder signs. Mice were monitored for symptoms of disease and euthanized as previously described
496 criteria for humane endpoint³⁹.

497

498 **Cryo-ET of LGTV-infected mouse brain tissue**

499 Based on optical projection tomography that visualized the infection distribution in entire, *ex vivo*
500 brains³⁹, choroid plexuses were surgically removed from brains of LGTV-infected *Ifnar*^{-/-} mice *post*
501 *mortem*. The choroid plexuses were perfused with ice-cold phosphate-buffered saline (PBS) and
502 rapidly transferred to ice-cold artificial CSF⁷⁶. Just prior to high-pressure freezing, the tissue was
503 placed in a 3 mm copper high-pressure freezing carrier (Wohlwend) which can be clipped into an
504 Autogrid. The sample was covered with a 20% dextran solution in PBS as cryoprotectant and covered
505 with a sapphire disk. The assembled carrier was rapidly vitrified using a Leica EM HPM100 high-
506 pressure freezer. The frozen carrier was trimmed at cryogenic temperatures using a Leica EM FC7
507 cryo-ultramicrotome with a diamond knife. The copper carrier was trimmed to leave a flat tissue
508 sample on the carrier, measuring 100 μm in width, 20 μm in thickness, and 30 μm in depth⁷⁷. Frozen
509 carriers were clipped into Autogrids (ThermoFisher) prior to cryo-FIB milling with a Scios dual-beam
510 FIB/SEM microscope (ThermoFisher Scientific).

511 The sample was coated with a protective platinum layer using a gas injection system for 15 seconds at
512 a working distance of 7 mm. The cryostage was tilted at an angle of approximately 10° for milling. A
513 rough milling was initially performed with an ion beam accelerating voltage of 30 kV and a current
514 ranging from 0.79 to 2.5 nA to reach a thickness of 1 μm . Additionally, the two sides of the lamella
515 were milled above and below to allow cryo-ET data collection by preventing the thick edges of the
516 tissue from obstructing transmission EM imaging⁷⁷. After rough milling, one edge of the lamella was
517 detached from the main platform to relieve stress. When the lamella reached a thickness of
518 approximately 1 μm , the ion beam current was lowered to 80 to 230 pA for fine milling, resulting in a
519 final lamella thickness of around 200 nm. SerialEM was used to collect tilt series data with tilt angles
520 ranging from 40° to -40° in 2° increments. The total electron dose for a single tilt series was
521 approximately 100 e-/ \AA^2 , with defocus between -5 and -10 μm . Tomograms were generated as
522 described above for cells.

523

524 **Statistics and reproducibility**

525 Data and statistical analysis were performed using Prism (GraphPad Software Inc., USA). Details
526 about replicates, statistical test used, exact values of n, what n represents, and dispersion and precision
527 measures used can be found in figures and corresponding figure legends. Values of $p < 0.05$ were
528 considered significant. All tomograms shown are representative of larger data sets as indicated in
529 Table 1.

530

531 **Data availability**

532 The cellular subtomogram averages of immature and mature Langat virus are deposited at the Electron
533 Microscopy Data Bank with accession codes EMD-51640 and EMD-51642, respectively.

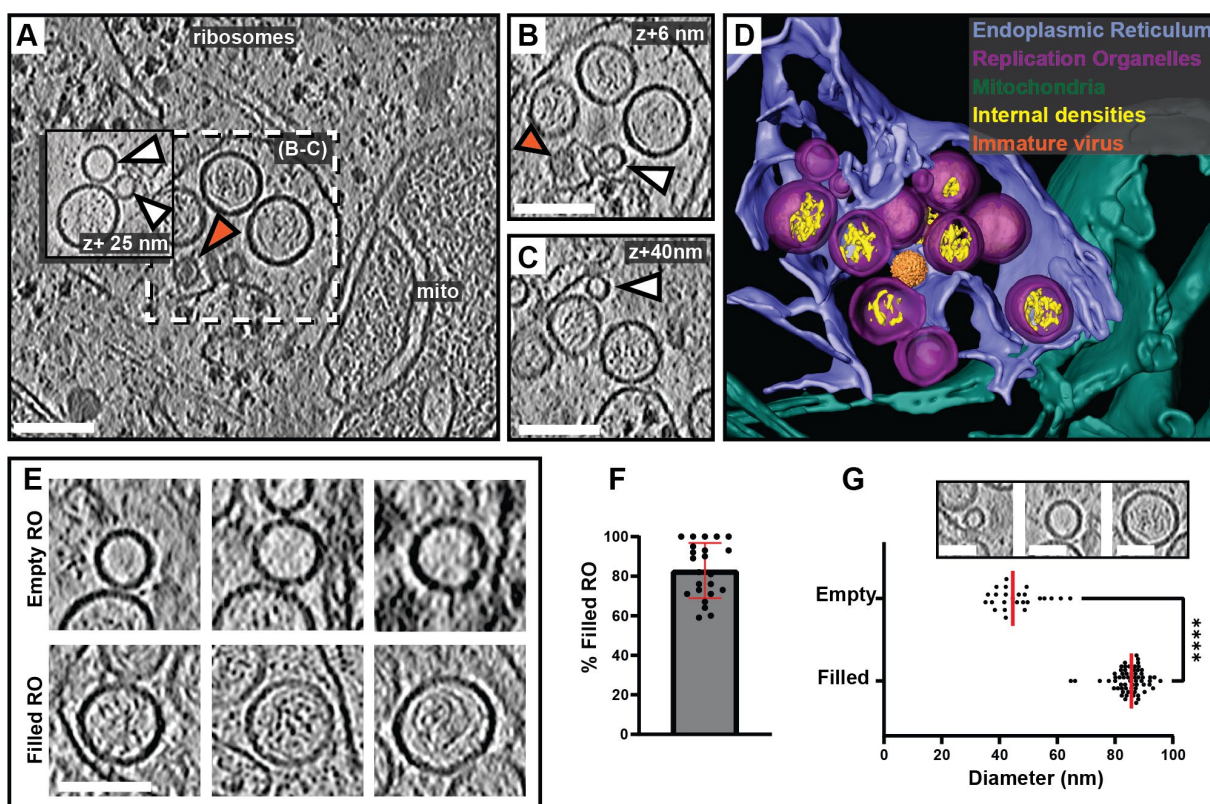
534

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547 performed at the Umeå Center for Electron Microscopy (UCEM) a SciLifeLab National Cryo-EM
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551 Renner for valuable comments and suggestions.

552

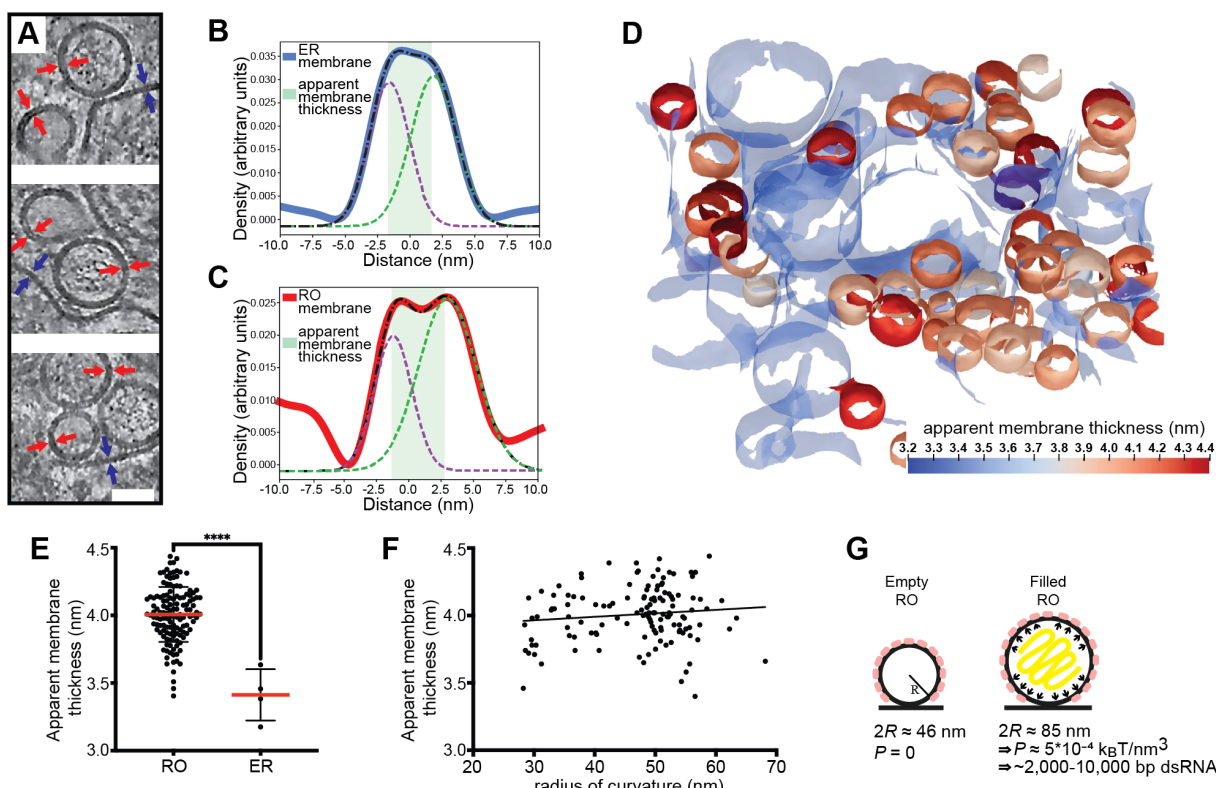
553 Figures and Tables



554

555 **Figure 1: *In situ* cryo-ET uncovers two states of Langat virus replication organelles.** (A) Slice
556 through a tomogram of FIB milled LGTV-infected cell showing viral ROs enclosed within the ER
557 with an immature virion (orange arrow). (B-C) Close up views of the outlined framed region in (A) at
558 their respective Z heights in the tomogram. (A-C) White arrows indicate empty ROs. (D)
559 Segmentation of the tomogram in (A), with color schemes defined for each structure. The immature
560 virion is represented by a subtomogram average. (E) Representative examples of empty and filled viral
561 RO observed in cryo-tomograms of milled LGTV-infected cells. (F) Percentage of filled ROs observed
562 in 23 tomograms of LGTV-infected cells. (G) Size distribution of empty (n=25) and filled (n=63) RO
563 observed in tomograms. The inset represents the different sizes of viral spherules observed in the
564 tomograms. (F-G) Red lines represent the average, each dot corresponds to one analyzed tomogram
565 (see also Table 1). Statistical significance by unpaired two-tailed Student's *t* test: ****p* < 0.0001.
566 Scale bars 100 nm.

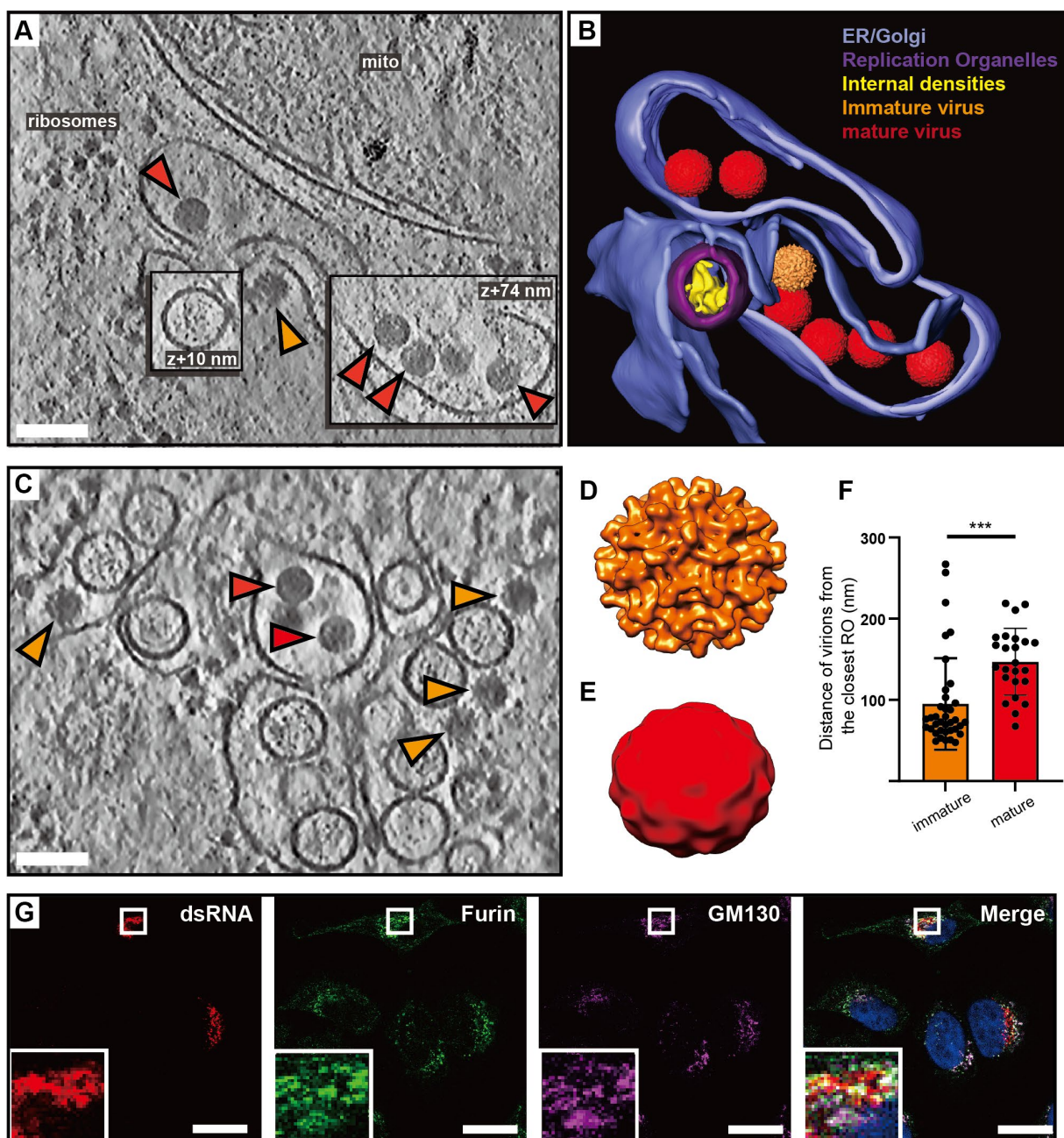
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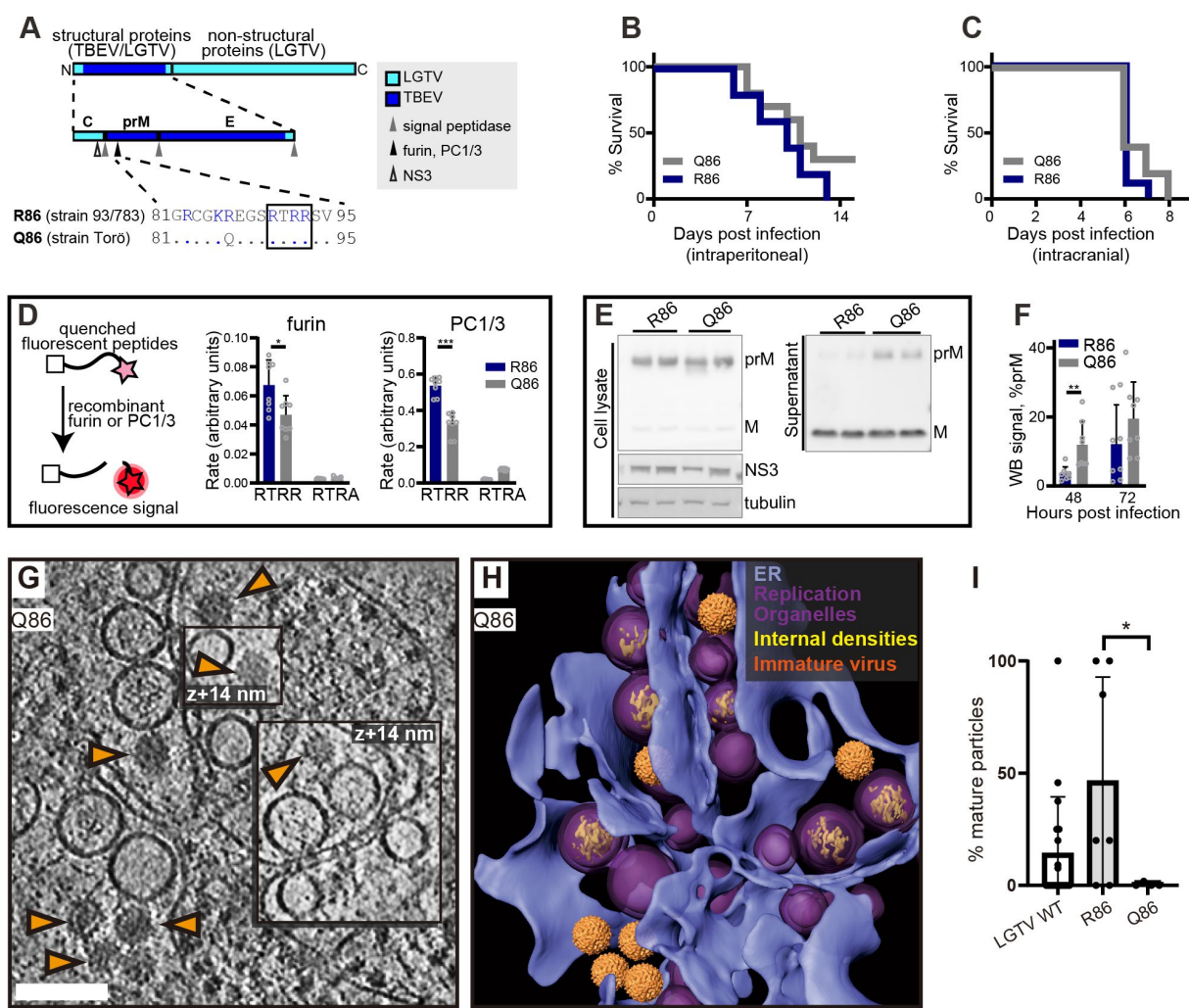
569 **Figure 2: The influence of a membrane coat and viral RNA in shaping the replication organelles.**
570 (A) Slices through tomograms of empty and filled ROs in LGTV-infected cells. Blue and red arrows
571 indicate the thickness of the ER and RO membranes, respectively. Scale bar, 50 nm. (B-C) Membrane
572 thickness estimation by dual Gaussian fitting to radial density plots through a representative ER
573 membrane (B) and RO membrane (C). Solid lines, membrane density profile; dashed lines, fitted
574 composite and dual Gaussian; shaded area, estimated thickness. (D) ER and ROs membranes from a
575 representative tomogram of an LGTV-infected cell, color coded by apparent membrane thickness. ER
576 membrane is partially transparent. (E) Apparent membrane thickness quantification in four tomograms
577 of LGTV-infected cells, comparing individual ROs ($n=132$) and the surrounding ER ($n=4$). Red lines,
578 average. Statistical significance by unpaired two-tailed Student's t test: *** $p < 0.0001$. (F)
579 Relationship between radius of curvature and apparent membrane thickness for individual ROs ($n=132$
580 from six tomograms). (G) Model of the mechanisms determining viral RO size. Two RO states exist in
581 infected cells: empty ROs with a baseline size set independent of luminal RNA, and filled ROs, whose
582 larger size is due to intraluminal pressure from $\sim 2,000\text{-}10,000$ bp dsRNA.

583



585 **Figure 3: Virions form and mature in the immediate vicinity of replication organelles.** (A) Slice
 586 through tomogram of LGTV-infected cell showing an immature virion budding across from a RO
 587 (orange arrow), and mature virions (red arrows) in adjacent membranes. (B) Segmentation of the
 588 tomogram in (A), with color labels defined for each structure. (C) Slice through tomogram of LGTV-
 589 infected showing mature (red arrows) and immature (orange arrows) LGTV virions observed near the
 590 viral RO. (D-E) Subtomogram averages of immature (D) and mature (E) LGTV from cellular
 591 tomograms. (F) The distance from immature ($n=37$) and mature ($n=24$) virions to the closest RO, from
 592 five tomograms. Statistical significance by unpaired two-tailed Student's t test: *** $p < 0.005$. (G)
 593 Representative immunofluorescence micrograph of dsRNA, furin and Golgi marker GM130 in LGTV-
 594 infected cells at 24 h p.i. Rightmost panel: merge including DAPI-staining of nuclei (blue). Scale bars
 595 100 nm (A,C), 5 μ m (G).

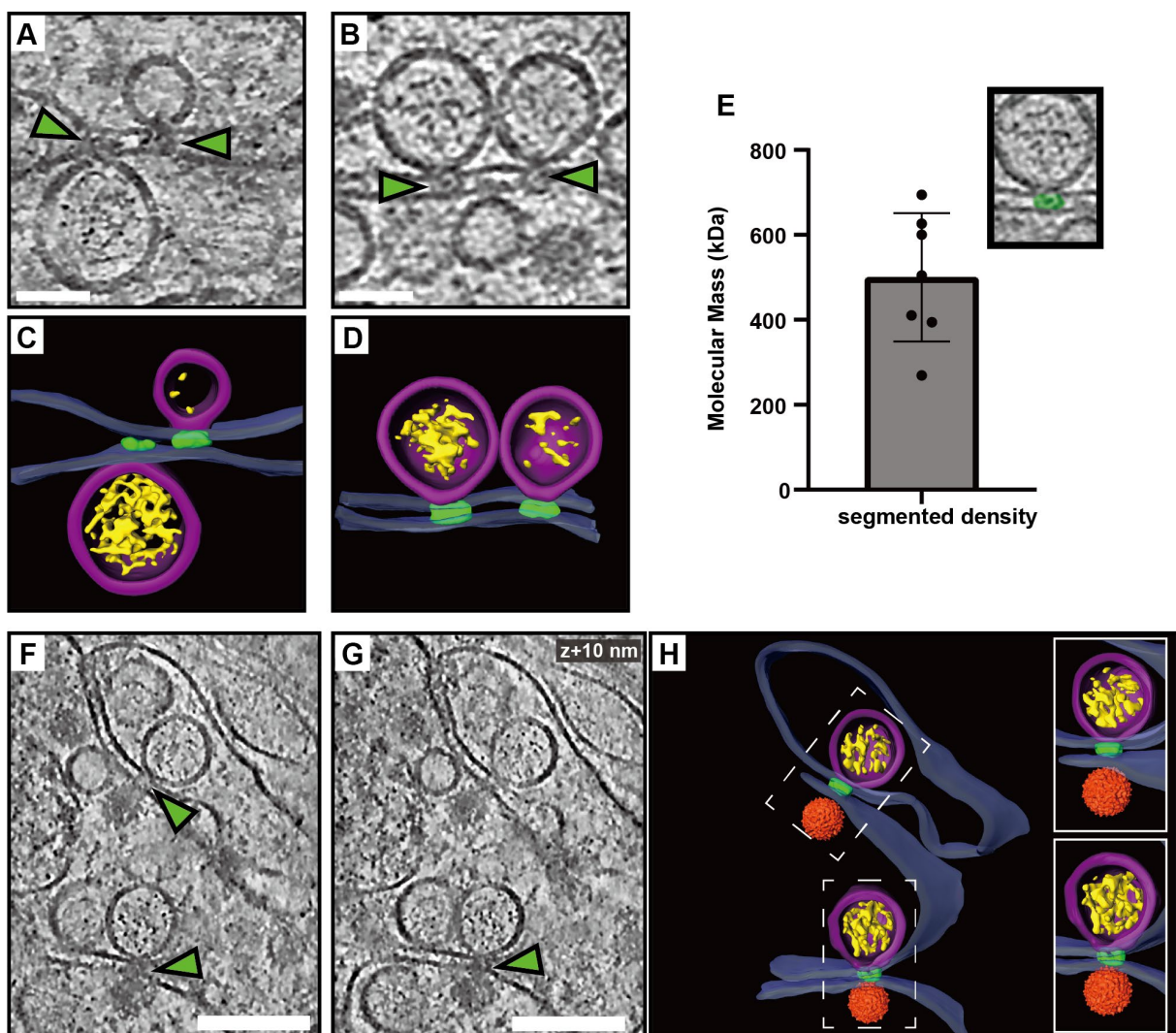
596



597

598 **Figure 4: The TBEV furin site variants R86 and Q86 differ in cleavage efficiency and replication**
 599 **organelle-proximal virion maturation.** (A) The polyprotein of a chimeric LGTV with prM and ecto-
 600 E from TBEV strain 93/783, as color coded. Recognition sites for viral and cellular proteases are
 601 shown within the structural protein region, and the furin site sequences from TBEV strains 93/783 and
 602 Torö are shown highlighting the difference at position 86 (., identical sequence). (B) Percentage
 603 survival of *Ips1^{-/-}* mice infected intraperitoneally with 10⁴ FFU with chimeric LGTV R86 (n=5) or Q86
 604 (n=10). (C) As (B), for but *Ips1^{-/-}* mice infected intracranially with 10² FFU (n=9, R86 and n=10,
 605 Q86). (D) Enzymatic cleavage using recombinant furin or PC1/3 with peptides corresponding to furin
 606 site sequences in (A) (RTRR), or peptides with impaired furin sites (RTRA). Data from four
 607 independent experiments performed in duplicates are shown, with mean values and standard deviation.
 608 (E) prM and M protein levels in cell lysates and supernatant 48 h.p.i. visualized by immunoblotting.
 609 Viral NS3 and cellular tubulin included as infection and loading control. Representative blots are
 610 shown. (F) Ratio of prM/M intensity quantified in supernatants at 48 and 72 h.p.i. Data from four
 611 independent experiments performed in duplicates are shown, with mean values and SD. (G) Slice
 612 through tomogram of chimeric LGTV Q86-infected cell showing a predominance of immature virions
 613 (orange arrows) within the cytoplasm. Scale bar 100 nm. (H) segmentation of the tomogram in (A),
 614 with color labels defined for each structure. (I) Percentage of mature virions in the tomograms of
 615 LGTV WT (n=19), chimeric LGTV R86 (n=6), LGTV Q86- infected cells (n=4). (D,F,I) Statistical
 616 significance by unpaired two-tailed Student's *t* test: *p<0.05, **p<0.01, ***p < 0.001.

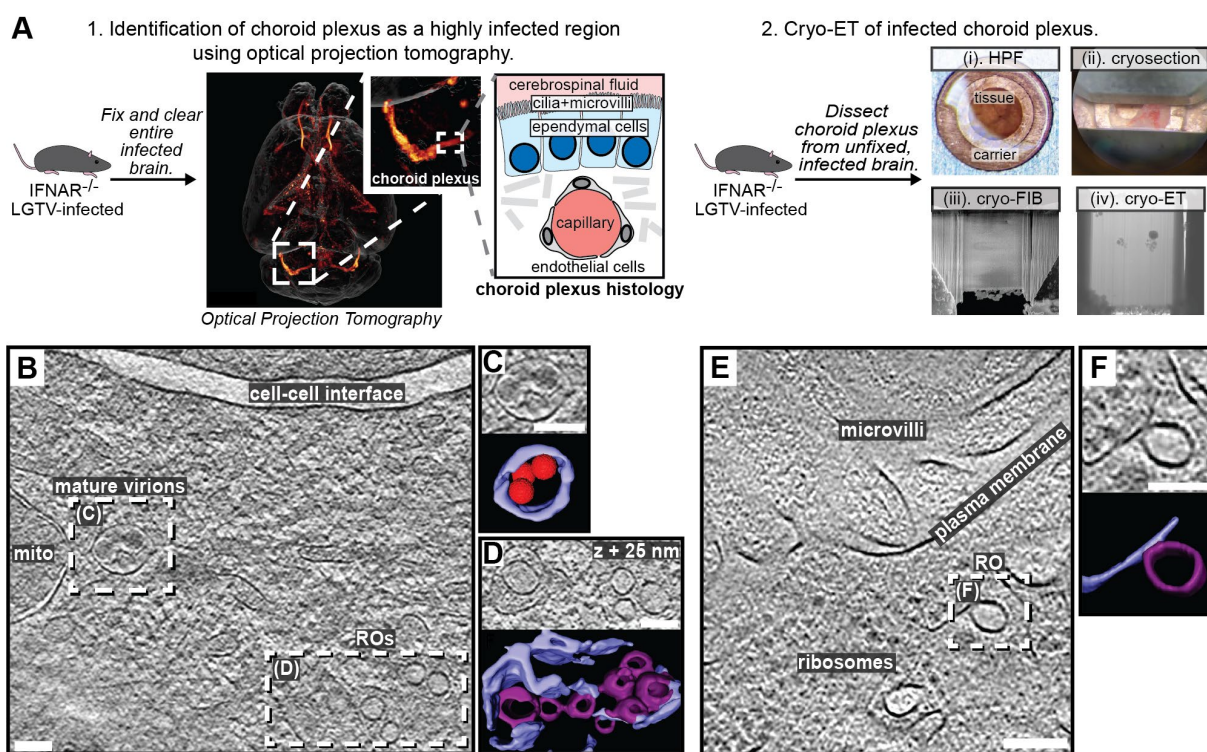
617



618

619 **Figure 5: A protein complex zippers replication organelles to an apposed ER membrane.** (A-B)
620 Slices through tomograms of LGTV-infected cells showing complexes (green arrows) located at the
621 neck of the ROs, connecting them to the adjacent ER membrane. (C-D) Segmentation of the
622 tomograms in (A-B). (E) Estimated molecular masses of the complex (n=7). (F-G) Slices through the
623 same tomogram at two different Z heights, showing complexes linking the RO to the site of virus
624 assembly (green arrows). (H) Segmentation of the tomogram shown in (F-G). (C-D,H) Blue, ER
625 membrane; purple, RO; yellow, luminal densities; green, neck complex; orange, immature virions.
626 Scale bars, 100 nm.

627



628

629 **Figure 6: LGTV replication visualized in *ex vivo* mouse brain using cryo-ET.** (A) Workflow for
630 cryo-ET of *ex vivo* mouse brain. LGTV-infected brains from *Ifnar*^{-/-} mice were processed differently
631 based on the imaging method. 1. Fixed and cleared brains were immunostained for the viral protein
632 NS5 (orange/red) and imaged using optical projection tomography. Zoomed insets show infected
633 regions of the choroid plexus, and its anatomy. 2. For cryo-ET, the choroid plexus from unfixed and
634 unstained LGTV-infected brains was (i) high-pressure frozen, (ii) trimmed using a cryo-
635 ultramicrotome, (iii) FIB milled, and (iv) transferred to a cryo-TEM for cryo-ET. (B) Slice through a
636 tomogram of LGTV-infected choroid plexus showing viral replication organelles and smooth virions
637 in proximity, enclosed within membrane vesicles. (C-D) Close-up of the areas indicated in (B) along
638 with their corresponding segmentations. (E) Slice through a tomogram of LGTV-infected choroid
639 plexus showing a *bona fide* RO with a thicker membrane than the ER, consistent with observations
640 from Fig. 2. (F) Close-up of the area indicated in (E) along with its corresponding segmentation. (B-F)
641 Colors as in Fig. 3. Scale bars, 100 nm.

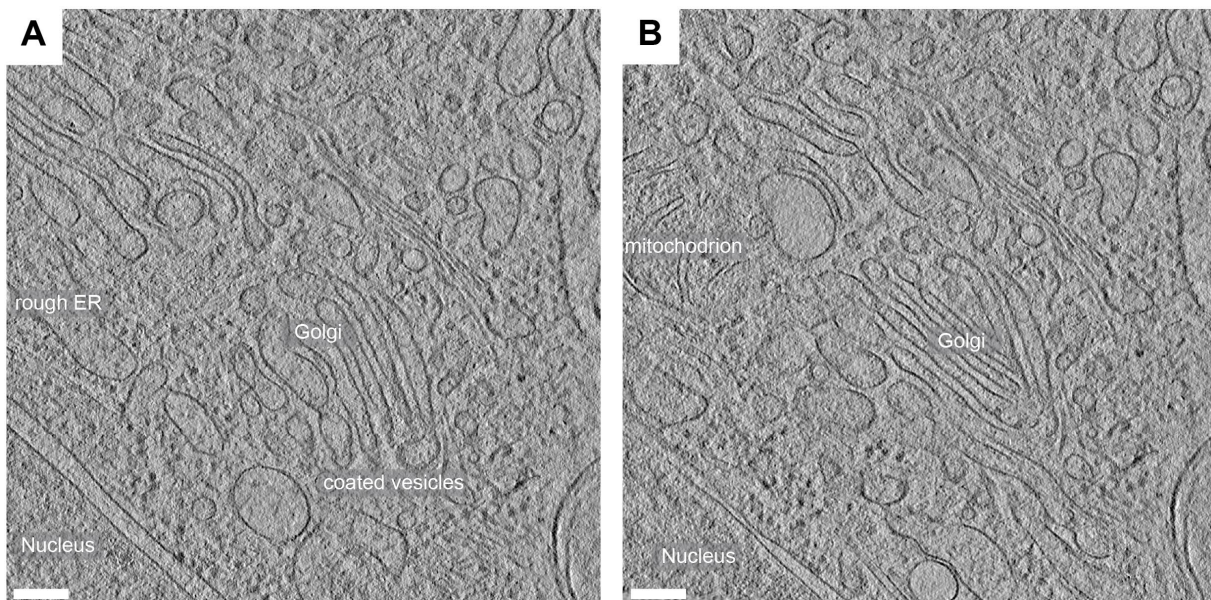
642

643 **Table 1. Number of tilt series recorded and tomograms analyzed.** “Number of tilt series” refers to
644 the total number of tilt series recorded on a given sample. All of these tilt series were used to
645 reconstruct tomograms, and the “number of tomograms with events” refers to the number of
646 tomograms with virus replication-related events.

sample	number of tilt series	number of tomograms with events
LGTV WT	51	23
rLGTV ^{T:prME} R86	12	7
rLGTV ^{T:prME} Q86	18	7
LGTV in <i>ex vivo</i> choroid plexus	45	8

647

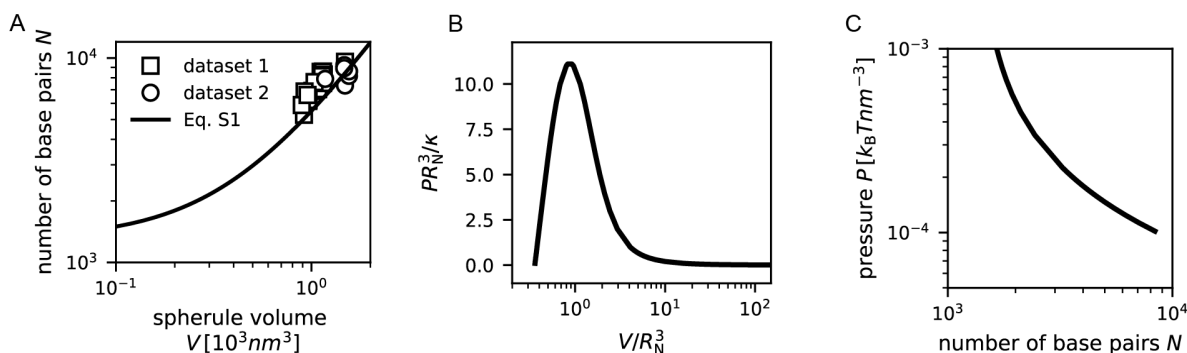
648 **Supplementary figures and movies**



649

650 **Figure S1: cryo-ET of uninfected A549 cells.** (A-B) Slices from two tomograms of uninfected A549
651 cells reveal typical cytoplasmic features, as indicated, including a non-dilated ER and *bona fide* Golgi
652 cisternae with typical morphology. Scale bars, 100 nm.

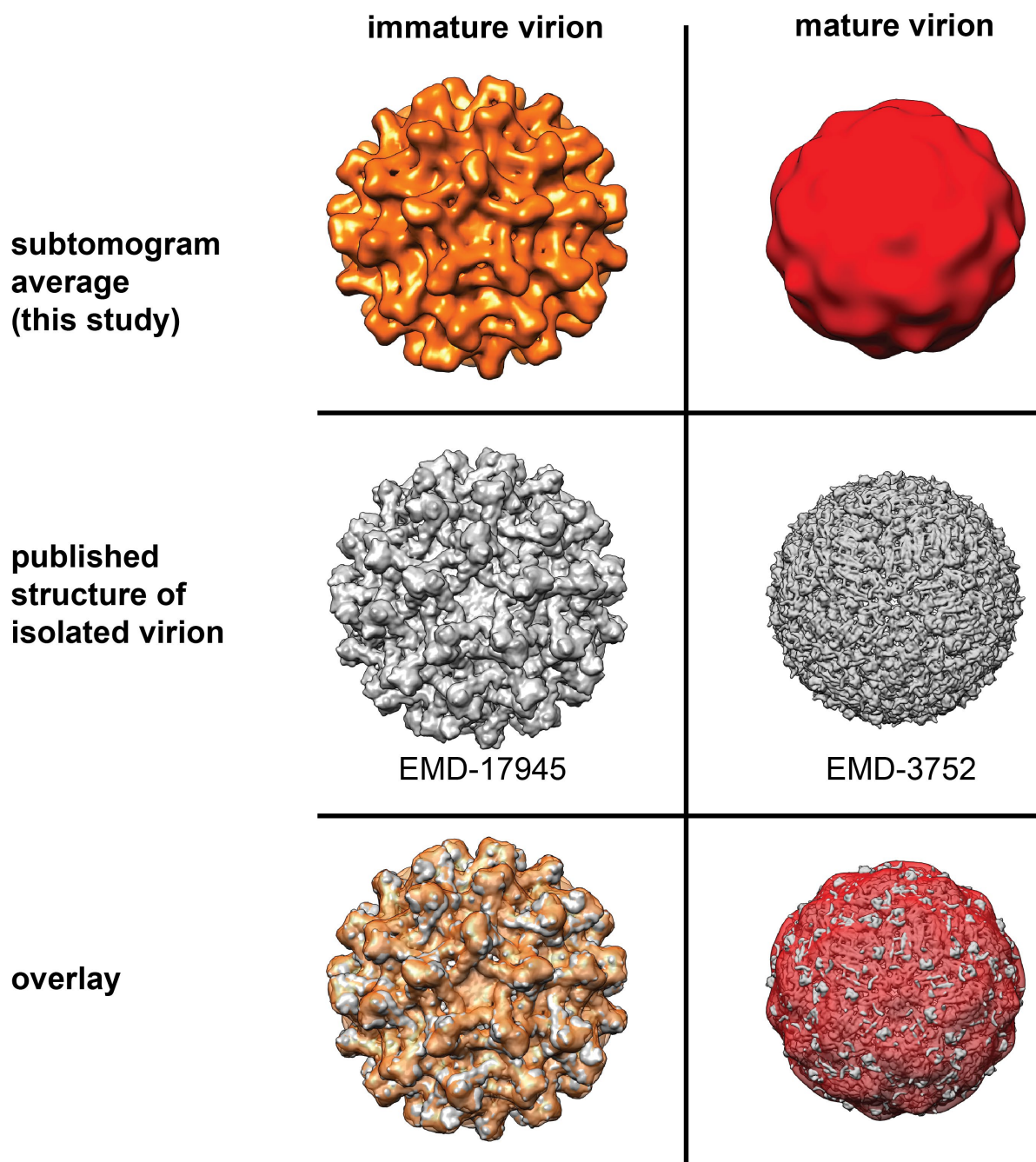
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654

655 **Figure S2. Pressure exerted by an RNA strand.** (A) Relation between number of RNA base pairs
656 and RO volume. The data is reproduced from Laurent *et al*³². We note that in Laurent *et al*³² the RNA
657 length is shown, while here the number of base pairs is shown, assuming an interbasepair distance of
658 2.56 Å. (B) Relation between the scaled pressure and the scaled volume. The details of the underlying
659 model are presented in Laurent *et al*³². (C) Relation between number of RNA base pairs N and
660 pressure P, where we use the results from (A-B) to convert the RO volume into number of RNA base
661 pairs.

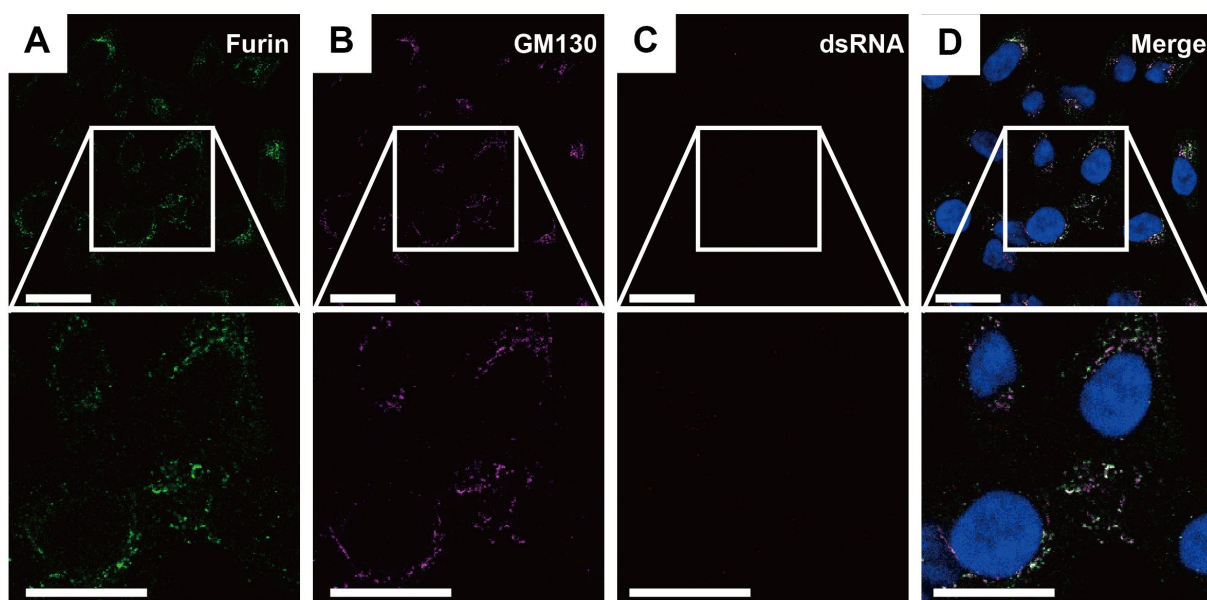
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663

664 **Figure S3: Comparison of cellular subtomogram averages with isolated virion structures.** The
665 cellular subtomogram averages from this study (top row) are compared to low-pass filtered published
666 structures of immature and mature TBEV, from Fuzik *et al*²⁵ and Anastasina *et al*²³, respectively (mid
667 row). The overlays (bottom row) were created using the Align to Volume command, and are shown
668 with the subtomogram averages in semi-transparent surface representation.

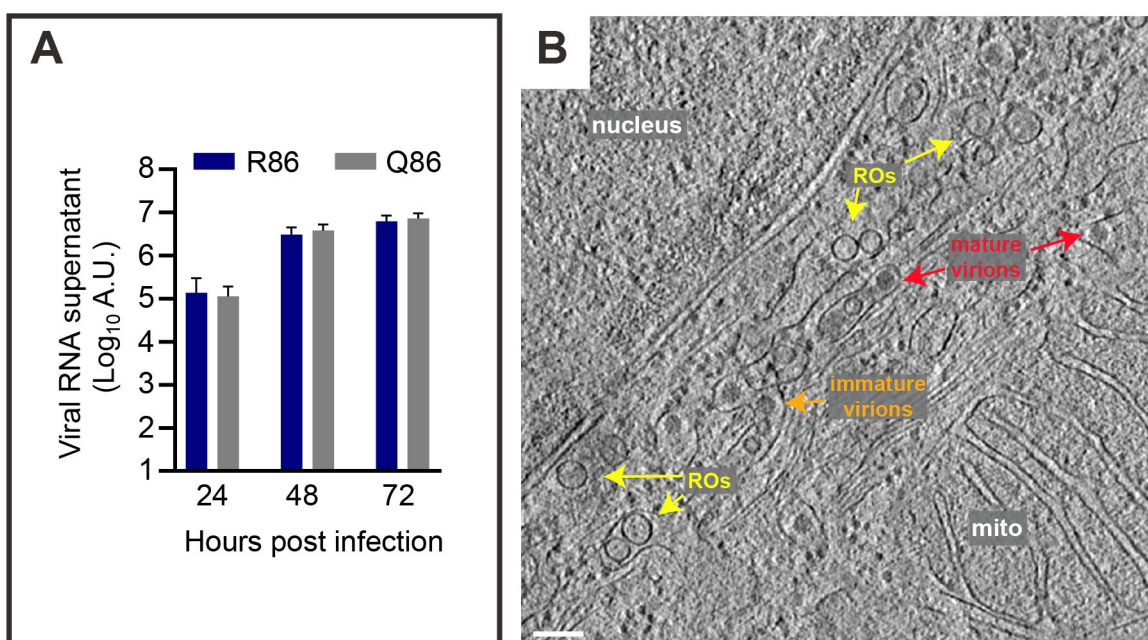
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670

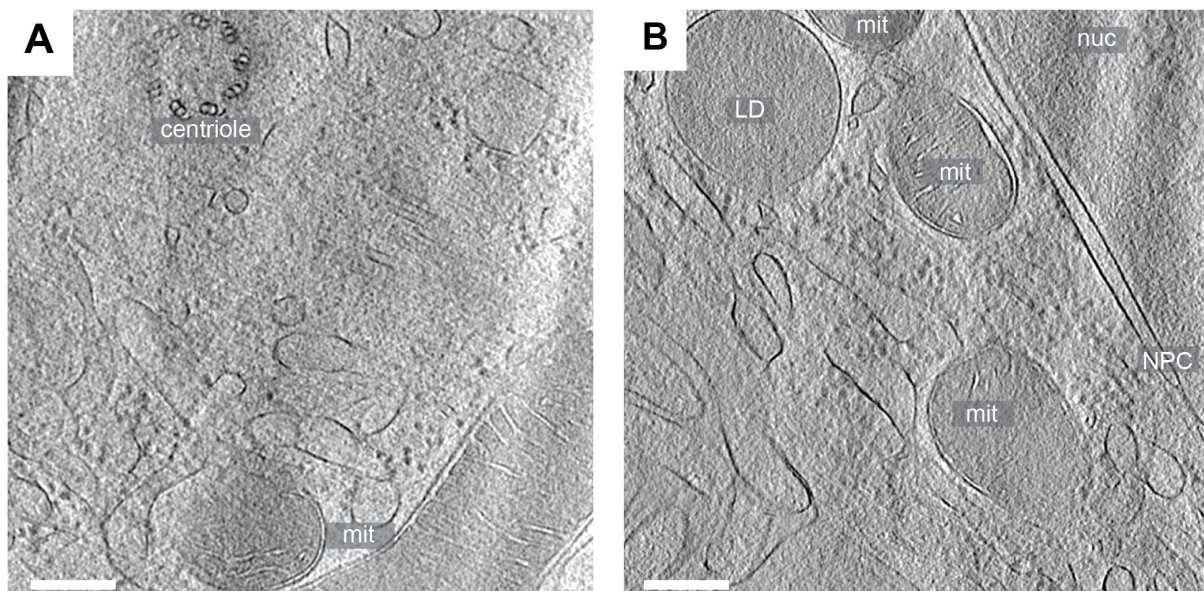
671 **Figure S4: Furin localization in uninfected cells.** Immunofluorescence microscopy of uninfected
672 cells showing furin (A) and Golgi marker GM130 (B) and their colocalization in the absence of viral
673 infection (D). Additional channels contain dsRNA staining (C) and DAPI staining of cell nuclei (D).
674 Scale bars, 10 μ m.

675



676

677 **Figure S5: Data on rLGTvT:prME Q86 and R86.** (A) Growth kinetics of rLGTvT:prME R86 and Q86
678 upon infection of A549 at MOI 1, quantitated as the amount of viral RNA in supernatant per qPCR.
679 (B) Slice from a tomogram of a cell infected with rLGTvT:prME R86 at 24 h p.i. showing various
680 cytoplasmic and virus-related features, as indicated. Scale bar 100 nm.



681

682 **Figure S6: Features unrelated to infection in cryo-electron tomograms of *ex vivo* brain tissue.** (A-
683 B) Slices from two tomograms of high-pressure frozen choroid plexus from LGTV-infected *Ifnar*^{-/-}
684 mice. The features indicated are a centriole, several mitochondria (mit), a lipid droplet (LD), the
685 peripheral area of a nucleus (nuc) and the nuclear envelope including one nuclear pore complex
686 (NPC). Scale bar, 100 nm.

687

688 **Movie S1.** Tomographic volume corresponding to the slice shown in Fig. 1A.

689 **Movie S2.** The segmentation shown in Fig. 1D.

690 **Movie S3.** Tomographic volume corresponding to the slice shown in Fig. 3A.

691 **Movie S4.** The segmentation shown in Fig. 3B.

692 **Movie S5.** Tomographic volume corresponding to the slice shown in Fig. 4G.

693 **Movie S6.** The segmentation shown in Fig. 4H.

694 **Movie S7.** Tomographic volume corresponding to the slices shown in Fig. 5F-G.

695 **Movie S8.** The segmentation shown in Fig. 5H.

696 **Movie S9.** Tomographic volume corresponding to the slice shown in Fig. 6B.

697

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