

1 *Ex vivo* midgut cultures of *Aedes aegypti* are efficiently infected by mosquito-borne 2 alpha- and flaviviruses

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24 **Abstract (250 words)**

25 *Aedes aegypti* mosquitoes can transmit several arboviruses, including chikungunya virus
26 (CHIKV), dengue virus (DENV), and Zika virus (ZIKV). When blood-feeding on a virus-
27 infected human, the mosquito ingests the virus into the midgut (stomach), where it
28 replicates and must overcome the midgut barrier to disseminate to other organs and
29 ultimately be transmitted via the saliva. Current tools to study mosquito-borne viruses
30 (MBVs) include 2D-cell culture systems and *in vivo* mosquito infection models, which
31 offer great advantages, yet have some limitations.

32 Here, we describe a long-term *ex vivo* culture of *Ae. aegypti* midguts. Cultured midguts
33 were metabolically active for 7 days in a 96-well plate at 28°C and were permissive to
34 ZIKV, DENV, Ross River virus (RRV) and CHIKV. *Ex vivo* midguts from *Culex pipiens*
35 mosquitoes were found to be permissive to Usutu virus (USUV). Immunofluorescence
36 staining confirmed viral protein synthesis in CHIKV-infected midguts of *Ae. aegypti*.
37 Furthermore, fluorescence microscopy revealed replication and spread of a reporter DENV
38 in specific regions of the midgut. In addition, two known antiviral molecules, β -D-N⁴-
39 hydroxycytidine (NHC) and 7-deaza-2'-C-methyladenosine (7DMA), were able to inhibit
40 CHIKV and ZIKV replication, respectively, in the *ex vivo* model.

41 Together, our results show that *ex vivo* midguts can be efficiently infected with mosquito-
42 borne alpha- and flaviviruses and employed to evaluate antiviral drugs. Furthermore, the
43 setup can be extended to other mosquito species. *Ex vivo* midgut cultures could thus be a
44 new model to study MBVs, offering the advantage of reduced biosafety measures
45 compared to infecting living mosquitoes.

46 **Importance (150 words)**

47 Mosquito-borne viruses (MBVs) are a significant global health threat since they can cause
48 severe diseases in humans, such as hemorrhagic fever, encephalitis, and chronic arthritis.
49 MBVs rely on the mosquito vector to infect new hosts and perpetuate virus transmission.
50 No therapeutics are currently available. The study of arbovirus infection in the mosquito
51 vector can greatly contribute to elucidating strategies for controlling arbovirus
52 transmission. This work investigated the infection of midguts from *Aedes aegypti*
53 mosquitoes in an *ex vivo* platform. We found several MBVs capable of replicating in the
54 midgut tissue, including viruses of major health importance, such as dengue, chikungunya,
55 and Zika viruses. Additionally, antiviral compounds reduced arbovirus infection in the
56 cultured midgut tissue. Overall, the midgut model emerges as a useful tool for diverse
57 applications such as studying tissue-specific responses to virus infection and screening
58 potential anti-arboviral molecules.

59 **1. Introduction**

60 Arboviruses are a diverse group of viruses that rely on arthropod vectors such as
61 mosquitoes, ticks, and sandflies to infect susceptible vertebrate hosts and perpetuate
62 transmission. Three arboviral families/orders are of most clinical relevance: *Bunyavirales*,
63 *Flaviviridae*, and *Togaviridae*, as they encompass important arboviruses of global health
64 concern, such as dengue virus (DENV), chikungunya virus (CHIKV), yellow fever virus,
65 and Zika virus (ZIKV) (1). These four mosquito-borne viruses (MBVs) are mainly
66 transmitted by female *Aedes* (Ae.) mosquitoes (e.g., *Ae. aegypti* and *Ae. albopictus*) (2),
67 with *Ae. aegypti* mosquitoes considered highly competent vectors because of their
68 anthropophilic behavior (3).

69 When blood-feeding on a virus-infected human or animal, a female mosquito will ingest
70 the virus-containing blood in the midgut (i.e., mosquito stomach), where the virus must
71 replicate and overcome the midgut barrier to subsequently disseminate to mosquito
72 secondary organs. As the mosquito becomes systemically infected, resulting in a high viral
73 load, the virus will reach the salivary glands. The virus will be released in the saliva upon
74 the next bite, allowing virus transmission when the mosquito feeds on uninfected hosts (4,
75 5). The midgut is thus the initial site for infection and replication of any arbovirus in the
76 mosquito. Virus replication in and spread across midgut cells are a requisite for a
77 productive arbovirus infection in the mosquito (4).

78 *In vitro* cell cultures can be useful to study arbovirus infection and other relevant processes
79 in mosquitoes; however, they present several limitations. Most of the established mosquito-
80 derived cell lines are not derived from tissues relevant to specific stages of mosquito-virus
81 interactions (e.g., salivary glands or midgut), but were generated from larval or embryonic
82 tissue (e.g., *C6/36* [larvae] and *Aag-2* [embryos] cells) (6). Therefore, results obtained from
83 *in vitro* experiments are ought to be taken cautiously. Furthermore, cell lines are considered
84 homogenous cultures, both genetically and phenotypically, and single-cell type
85 populations might not well represent multicellular tissues such as a midgut (7, 8).
86 Moreover, some biological differences might arise among the same cell line coming from
87 different laboratories (7).

88 Another method commonly used to study mosquito-virus interactions is the mosquito
89 infection model. Through this approach, mosquito infection occurs artificially via a
90 bloodmeal (membrane filled with infectious blood), and mosquitoes are sacrificed to assess
91 viral infection and vector competence (9). Although living-infected mosquitoes provide

92 valuable transmission data, the value of these data is directly proportional to the labor-
93 intensive nature of the method. Transmission studies require skilled personnel for the
94 handling of infected mosquitoes and facilities with a high biosafety level that ensure proper
95 containment, and carry the risks of prick injury with virus-infected dissection tools or of
96 infected mosquitoes escaping. Additionally, considerable variability among mosquito
97 infection studies might exist as colonies bearing the same name can be highly genetically
98 divergent among diverse laboratories (10).

99 *Ex vivo* organ cultures have been arising as an advantageous tool in research and have also
100 been described for insects. For instance, *ex vivo* culture of insect organs has been
101 successfully and widely defined for ixodid ticks. The synganglion, midgut, and salivary
102 glands dissected from ticks were viable for 10 days and permissive to both Langat and
103 Powassan viruses. (8, 11). On the contrary, reports on *ex vivo* organ cultures from mosquito
104 tissues are scarce and mainly focused on germline and fat body tissues (12, 13). Despite
105 the essential implication of the mosquito midgut in arbovirus replication and dissemination,
106 no *ex vivo* mosquito midgut culture has been properly described. To date, only a short-term
107 *ex vivo* assay in mosquito midguts has been reported, in which the midguts were only
108 maintained for 35 hours post dissection in tissue culture plates (14).

109 Nevertheless, *ex vivo* cultures could offer a unique perspective to (i) study vector-virus
110 interactions in a relevant tissue, for example, the mosquito midgut, (ii) reveal tissue-
111 specific responses to virus infection, or (iii) study tissue-specific metabolic processes.
112 Furthermore, *ex vivo* mosquito midgut cultures could provide a convenient and relevant
113 platform to generate results that can be extrapolated to living mosquitoes. Additionally, *ex*
114 *vivo* midgut cultures provide a controlled environment in which many parameters can be

115 easily adjusted, and they constitute a safer option due to the low degree of containment
116 required while working with BSL2/3 arboviruses, compared to working with live mosquito
117 infection models.

118 On the lookout for novel tools that could be of use to study arbovirus infection within the
119 arthropod vector, we dissected *Ae. aegypti* midguts and established their viability for a 7-
120 day period. We also infected the midgut tissues with several mosquito-borne alpha- and
121 flaviviruses, confirming the production of infectious viral particles. Furthermore,
122 previously reported antiviral drugs were able to reduce arbovirus replication in the treated
123 midguts. We thus developed an *ex vivo* mosquito midgut model that constitutes a valuable
124 tool to study MBVs which complements the arsenal available to the mosquito arbovirus
125 research field.

126 **2. Results**

127 **Viability of *Ae. aegypti* midguts cultured *ex vivo***

128 We first determined the viability of the dissected mosquito midguts in 96-well culture
129 plates. Since motility could be observed along the gut tissue following the dissection, a
130 bioassay was performed based on the number of contractions by the hindgut
131 (Supplementary material, Movie 1). The same area of the hindgut was analyzed for all
132 biological replicates (Fig. 1, b). Dissected midguts presented peristalsis in the hindgut for
133 up to 7 days post dissection (d.p.d.) (Fig. 1, d), indicating that the tissue was viable. At 10
134 d.p.d., the mean frequency in the *ex vivo* cultured midguts was significantly reduced
135 compared to day 0 measurements (day 0: 14.37 s vs day 10: 6.74 s) (Fig. 1, d), suggesting
136 that the tissue's viability had decreased.

137 We further assessed the metabolic activity of the *ex vivo* cultured midguts in a resazurin
138 salt-based assay. The dissected midguts remained metabolically active up until 7 d.p.d.
139 (Fig. 1, e), corresponding with the data obtained from gut peristalsis. Fluorescence intensity
140 in the tested midguts was still slightly higher than the “no organ” controls (NO, used to
141 measure background signal) and negative controls (NC, paraformaldehyde fixated
142 midguts) at 8, 9, and 10 d.p.d.. However, there was more variability among biological
143 replicates and therefore 7 d.p.d. was selected as the end time point for following
144 experiments.

145 Additionally, the effect of the neurotransmitter serotonin and the anticholinergic drug
146 atropine on peristalsis was assessed to rule out that this visual feature might be a reflex.
147 Midguts (day 0 post dissection) were exposed to either serotonin, atropine, or midgut
148 medium (mock-exposure), after which their gut peristalsis was recorded. The mean
149 frequency displayed by the mock-exposed midguts was 7.349 s (5 biological replicates,
150 median value: 7.48 s). Midguts exposed for 1 hour to serotonin displayed a slight increase
151 in gut motility, with a mean peristaltic period of 12.76 s (6 biological replicates, median
152 value: 10.61 s), whereas atropine-exposed midguts exhibit a somewhat reduced peristalsis,
153 with a mean peristaltic period of 6.72 s (6 biological replicates, median value: 7.58 s).
154 Despite the observed trends, there were no statistically significant differences among the
155 conditions evaluated (Supplementary Fig. S1).

156 **Infection of dissected *Ae. aegypti* midguts with mosquito-borne alpha- and
157 flaviviruses**

158 Next, *ex vivo* virus infection was performed in the cultured midguts to test their capacity
159 to support the replication of several arboviruses. Replication kinetics were evaluated for

160 Ross River virus (RRV) and CHIKV as representative members of the alphavirus genus of
161 the *Togaviridae* family. Both viruses were able to efficiently replicate in the *ex vivo*
162 cultured midguts with an increase in viral RNA levels between 1 and 3 days post infection
163 (d.p.i.), compared to viral RNA levels at 2 hours p.i. (h.p.i.) (Fig. 2, a). Infectious virus
164 titers of RRV and CHIKV increased accordingly in the midgut tissue (Fig 2, b), while no
165 infectious virus was observed yet at 2 h.p.i. (data not shown). The peak of viral RNA and
166 viral titer levels was detected at 3 d.p.i. for both alphaviruses.

167 To evaluate whether *Aedes*-borne flaviviruses could replicate in the *ex vivo* midguts, the
168 replication kinetics of DENV-2 and ZIKV were studied. As both viruses needed a longer
169 period to replicate in the midgut tissue (data not shown), viral RNA and infectious virus
170 levels were measured starting at 5 d.p.i.. Viral titers were increased at 5 d.p.i. and further
171 peaked at 7 d.p.i., with DENV-2 reaching RNA levels up to 2.5×10^8 genome
172 copies/midgut and a viral titer of 21 PFU/midgut. On the contrary, ZIKV replication and
173 infection presented more variability among individual midguts. The maximum amount of
174 ZIKV RNA and infectious virus quantified at 7 d.p.i. were 2.1×10^6 genome copies/midgut
175 and 4.2×10^2 PFU/midgut, respectively (Fig 3).

176 **Usutu virus (USUV) can modestly replicate in *ex vivo* cultured midguts from *Culex*
177 *pipiens* mosquitoes**

178 As a proof-of-concept, the *ex vivo* midgut model was applied to another mosquito species:
179 *Culex (Cx.) pipiens*. The midguts of *Cx. pipiens* thrived in culture and showed peristaltic
180 movements during their incubation, as *Ae. aegypti* midguts did. Therefore, we evaluated
181 the susceptibility of these *Cx.* midguts to USUV, as *Cx. pipiens* mosquitoes are considered
182 a competent vector for this virus (15). USUV infection was assessed in the midguts at 7

183 d.p.i. by qRT-PCR and plaque assay. Only 50% (3 out of 6) of the midguts became infected
184 with USUV, reaching up to 1.8×10^6 genome copies/midgut. However, only 2 out of these
185 3 USUV-infected midguts contained infectious virus, amounting to 6.4×10^2 PFU/mL. Of
186 note, a considerable amount of viral RNA was detected in samples corresponding to 2 h.p.i.
187 and in 2 out of the 3 negative controls included in the assay, yet no infectious virus was
188 detected in these samples (Fig. 4, a, b).

189 **CHIKV viral protein synthesis in infected *ex vivo* midguts**

190 Previous infection experiments indicated the susceptibility of the *ex vivo* midguts to
191 infection with several arboviruses. To further corroborate these results, the CHIKV E2
192 glycoprotein was visualized in CHIKV-infected midguts at day 3 p.i. by immunostaining.
193 Specific staining for this viral protein could be observed in the infected guts, while no
194 signal was observed in the negative controls (fixated guts that followed the same infection
195 protocol; Fig. 5, a). At day 3 p.i., E2 protein synthesis was mainly detected and spread
196 along the posterior midgut region (Fig. 5, b, c), with some infection foci located in the
197 hindgut region (Fig. 5, c). The E2 signal was also present in the tracheal tubes that remained
198 attached to the midgut after dissection (Fig. 5, b, arrow heads).

199 **Infection of *ex vivo* cultured midguts with an mCherry-expressing DENV-2**

200 To follow the progression of virus infection in the midguts by imaging, cultured mosquito
201 midguts were infected with DENV-2 expressing the red fluorescent protein mCherry
202 (DV2/mCherry,(16)) and imaged using the FLoid™ Cell Imaging Station (Life
203 Technologies) at selected time points p.i.. The mCherry signal was used as a proxy for
204 infection. DENV-2 mCherry infection was observed initially at day 3 p.i. as single or few
205 foci in the posterior midgut region (Supplementary Fig. S2, a). Over time, these focal

206 infection points increased in number and spread to neighboring areas, primarily along the
207 posterior region of the midgut and the hindgut (Supplementary Fig. S2, b-e). No mCherry
208 expression was observed in the negative control or mock-infected midguts at day 7 p.i.
209 (Fig. 6, a). Following 7 days of infection, the mCherry signal had spread mostly along the
210 posterior midgut region, forming infection foci comprising multiple cells (Fig. 6, b, c).
211 Moreover, some infected cells were found in the tracheal tubes that remained in the midgut
212 tissue after dissection (Supplementary Fig. S2, h).

213 **Arbovirus replication is reduced upon treatment with antiviral drugs in midgut
214 cultures**

215 We next assessed whether the *ex vivo* mosquito midguts could be used to evaluate the
216 antiviral activity of inhibitors against arboviruses. To this end, the antiviral activity of β -
217 D-N⁴-hydroxycytidine, also known as EIDD-1931 or NHC, was tested against CHIKV at a
218 concentration of 50 μ M. At day 3 p.i., no difference in the CHIKV RNA levels was
219 observed between the untreated and NHC-treated groups (Fig. 7, a). In contrast, NHC
220 significantly reduced the infectious virus levels by 1 log (mean values, control: 1.0×10^4 ;
221 50 μ M NHC: 1.1×10^3 PFU/midgut) (Fig. 7, b).

222 As the *ex vivo* cultured midguts also supported flavivirus infection, the viral polymerase
223 inhibitor 7-Deaza-2'-C-Methyladenosine (7-DMA) was tested against ZIKV. Viral RNA
224 levels were reduced for all treated midguts (25, 100, and 200 μ M of 7-DMA) compared to
225 the untreated group at day 7.p.i.. However, only the groups treated with 7-DMA at 100 and
226 200 μ M showed a significant difference compared to the untreated group, with 1.5 log and
227 2.4 log reductions (virus control: 2.6×10^6 ; 100 μ M: 7.0×10^4 ; 200 μ M: 9.5×10^3 mean
228 genome copies/midgut), respectively (Fig. 7, c).

229 **3. Discussion**

230 The study of arbovirus infection in the mosquito vector relies mainly on a combination of
231 *in vitro* and *in vivo* approaches, which has yielded great progress in the knowledge
232 regarding arbovirus biology, virus-vector interactions, and vector competence. While
233 convenient and handy, *in vitro* cell culture systems have the shortcoming of not being as
234 biologically relevant as mosquito infection models, aside from the limited selection of
235 mosquito cell lines available. On the other hand, working with living infected mosquitoes
236 grants useful information when studying virus-vector interactions or vector competence for
237 a specific virus, but such *in vivo* models are not always accessible and require the
238 implementation of cumbersome safety measures ensuring adequate containment while
239 working with BSL2/3 pathogens (17). *Ex vivo* organ culture methods have not yet been
240 described in-depth for the study of MBVs. Hence, in this study, we have established an *ex*
241 *vivo* mosquito midgut model using dissected midguts from *Ae. aegypti* mosquitoes. When
242 cultured *ex vivo*, the midguts displayed peristaltic contractions, mainly observed as waves
243 along the hindgut, which is a frequent occurrence for this type of tissue (18). Here, we used
244 this visual feature as a proxy for viability, paired with a custom script for gut motility in
245 zebrafish (19) to measure the peristaltic period (time between contractions) in the *ex vivo*
246 cultured guts, which remained stable for 7 days p.d.. This observation was further
247 corroborated when measuring the metabolic activity of the dissected midguts over time in
248 culture with PrestoBlue™ (PB). This resazurin-based reagent has been successfully used
249 to assess cell viability and cytotoxicity in both two-dimensional cell monolayers and 3D
250 cellular interfaces (including organ explants) (20–23). Here, we found that only after 7

251 hours of incubation of the *ex vivo* midguts (using multiple midgut organs per well) in
252 presence of the PB reagent yielded a colorimetric change.

253 The *ex vivo* cultured mosquito midguts supported infection with four arboviruses, despite
254 the unusual route of infection (through incubation with virus inoculum) that differs from
255 what occurs naturally in mosquitoes (infection via blood feeding). Both viral RNA and
256 infectious virus particles were detected in the mosquito midgut organs following a
257 progressive increase over time, which also confirmed the viability of the midguts. These
258 replication kinetics results indicated the preservation of midgut tissue in an *ex vivo* culture
259 set-up and, more specifically, the presence of midgut cells that could constitute the target
260 for arbovirus infection.

261 With *Ae. aegypti* being the main vector for both DENV-2 and ZIKV, it was however
262 unexpected that infectious virus titers for both flaviviruses detected in the *ex vivo* midguts
263 did not reach higher levels than what was inoculated (1×10^4 PFU/mL), in contrast to viral
264 RNA levels. Kinetics of DENV replication in mosquitoes have reported a steady increase
265 in infectious virus until 8 d.p.i., after which it normally declined without affecting viral
266 RNA levels (24, 25). Such results could explain the discrepancy seen between flavivirus
267 RNA and infectious levels. More importantly, it has been described that within 2 hours of
268 infection with DENV or ZIKV (both *in vivo* and *ex vivo*), there is a rapid induction of
269 apoptosis in the *Ae. aegypti* midgut epithelium in an attempt of the host to control flavivirus
270 infection (26). Nonetheless, this process might cause tissue damage, which correlates with
271 the high midgut cell turnover rate in mosquitoes during a bloodmeal digestion (27). As
272 such, we cannot disregard that the *ex vivo* midguts in our experiments undergo the same
273 process during DENV or ZIKV infection, consequently damaging the tissue and thus

274 limiting flaviviral replication. This hypothesis may also apply to *ex vivo* midguts of *Cx.*
275 *pipiens* mosquitoes, as a comparable replication pattern can be observed when assessing
276 USUV (flavivirus) infection at day 7 p.i.. Further addition of an apoptosis inhibitor to the
277 virus inoculum used to infect the *ex vivo* midguts might elucidate whether rapid induction
278 of apoptosis in the midgut epithelium is indeed a limiting factor for flavivirus replication
279 in this setup.

280 Active virus replication was confirmed for CHIKV through an immunofluorescence assay.
281 Envelope (E2) protein synthesis was localized in the posterior midgut region, where it
282 would also occur when the mosquito ingests a CHIKV-infectious bloodmeal (28),
283 regardless of the infection method employed with the *ex vivo* cultured midguts. Moreover,
284 CHIKV infection of tracheal tubes in the tissue could be observed, consistent with *in vivo*
285 reports (28). Similar results were obtained when analyzing the progression of
286 DENV2/mCherry infection in the *ex vivo* midguts. Previously, DENV-2 infection in *Ae.*
287 *aegypti* (Chetumal strain) midguts was reported to start with individual infected epithelial
288 cells detected as early as 2 d.p.i., slowly progressing to infection foci consisting of multiple
289 cells until the whole midgut organ was infected at 7-10 d.p.i. (24). In agreement with this,
290 a less strong but akin infection pattern was observed in DENV2/mCherry-infected midguts,
291 with few foci of several infected cells by 7 d.p.i.. Together, these data show that arbovirus
292 infection occurred at considerable levels in *ex vivo* cultured midguts and therefore it could
293 be used to study other aspects of arbovirus infection and facilitate the collection of
294 preliminary data before experimenting with mosquitoes *in vivo*.
295 The use of antiviral compounds to inhibit virus infection in the mosquito vector is an
296 innovative concept that might reduce or block arbovirus transmission from mosquitoes to

297 humans. This idea involves antiviral molecules being ingested by adult mosquitoes when
298 they take a bloodmeal on a mammalian host undergoing antiviral treatment. As the
299 mosquito midgut is the entry point and key replication site for MBVs, studying the effect
300 of antiviral compounds in the midgut tissue would be of great interest. For this purpose,
301 the antiviral activity of two inhibitors was assessed in the *ex vivo* mosquito midguts. NHC
302 is a nucleoside analog that has been characterized as a potent antiviral drug against
303 alphaviruses *in vitro*, including CHIKV and Venezuelan equine encephalitis virus (VEEV)
304 (29, 30). Various assays point to the compound acting as a pyrimidine analog that may
305 target the viral polymerase domain of nsP4, provoking chain-termination (31). In addition,
306 NHC also induces a high level of mutations in virus-specific RNAs, resulting in lethal
307 mutagenesis (32). Consistent with these findings, CHIKV RNA levels were not reduced in
308 NHC-treated midguts, but there was a marked decrease in virus infectivity. The viral
309 polymerase inhibitor 7DMA has shown potent anti-ZIKV activity *in vitro* and delayed
310 disease progression in mice (33). In the *ex vivo* mosquito midguts, 7DMA significantly
311 reduced ZIKV RNA loads. Although the RNA levels presented some variability among the
312 ZIKV-infected midguts, the inhibitory effect of 7DMA in the *ex vivo* mosquito midguts
313 was significant and followed a dose-response relationship. These results indicate that *ex*
314 *vivo* midgut cultures could be used to rationally select potential arbovirus-blocking
315 molecules to be tested in living mosquitoes at a later stage.

316 In summary, we have established a long-term *ex vivo* mosquito midgut culture. To support
317 this model, we have (a) assessed the viability of the *ex vivo* cultured midguts, (b)
318 determined the replication kinetics of two alphaviruses (RRV and CHIKV) and two
319 flaviviruses (DENV-2 and ZIKV), (c) detected viral protein synthesis and followed live

320 arbovirus infection in infected midguts, (d) demonstrated that the *ex vivo* protocol can be
321 translated to other mosquito genera, and, lastly, (e) evaluated the antiviral activity of two
322 inhibitors against CHIKV and ZIKV. Altogether, we have provided a reference for the use
323 of *ex vivo* mosquito midguts as a tool to study arbovirus infection and related processes,
324 while offering groundwork for developing other *ex vivo* mosquito organ cultures that can
325 potentially provide insightful data, like salivary glands.

326 **4. Materials and Methods**

327 **4.1. Cells**

328 ***Mammalian cells***

329 African green monkey kidney cells (Vero cells, ATCC CCL-81) and Vero E6 cells (ATCC
330 CRL-1586) were cultured in minimum essential medium (MEM 1X) enriched with 10%
331 Fetal Bovine Serum (FBS), 1% sodium bicarbonate, 1% L-glutamine and 1% non-essential
332 amino acids (NEAA). Baby hamster kidney cells (BHK, ATCC CCL-10) were maintained
333 in Dulbecco's Modified Eagle's Medium (DMEM) containing 10% FBS, 1% sodium
334 bicarbonate and 1% L-glutamine. Mammalian cell cultures were incubated at 37°C, with
335 5% CO₂.

336 ***Mosquito cells***

337 *Ae. albopictus* larval cells (C6/36, obtained from ATCC, CRL-1660) were maintained in
338 Leibovitz's L-15 medium containing 10% FBS, 1% Penicillin-Streptomycin (PenStrep),
339 1% NEAA, and 1% HEPES buffer. Mosquito-derived cell lines were incubated at 28°C,
340 without CO₂.
341 For cell culture assays containing virus or virus-infected material, the concentration of FBS
342 in the medium was reduced to 2%, for both mammalian and mosquito cells. All cell culture

343 media and supplements were obtained from GibcoTM, ThermoFisher Scientific (Aalst,
344 Belgium).

345 **4.2. Viruses**

346 *Flaviviruses*

347 Dengue virus serotype 2 (DENV-2/TH/1974, isolated in 1974 from human serum collected
348 in Bangkok, Thailand, GenBank MK268692.1) was kindly provided by Prof. A. Failloux
349 (Institut Pasteur, Paris, France) (34); ZIKV (SL1602, Suriname strain, GenBank
350 KY348640.1) was acquired via the EVAg consortium (<https://www.european-virus-archive.com>). The infectious clone DENV-2 pDVWS601 used for the construction of the
351 DENV-2 reporter virus expressing the red fluorescent protein mCherry (DV2/mCherry,
352 New Guinea C strain, NGC, GenBank AF038403.1) was kindly provided by Prof. Andrew
353 Davidson (University of Bristol, Bristol, UK) (16).

355 *Alphaviruses*

356 Ross River virus (RRV) was received from the National Collection of Pathogenic Viruses
357 (UK; catalog number 0005281v); and CHIKV (Indian Ocean strain 899, GenBank
358 FJ959103.1) was generously provided by Prof. Drosten (University of Bonn, Bonn,
359 Germany) (35).

360 Virus stocks were prepared by passaging the isolates on Vero (for CHIKV and USUV) or
361 C6/36 cells (for ZIKV, DENV-2, and RRV). Viral titers of the stocks were determined via
362 plaque assay or end-point titration on Vero or BHK cells.

363 **4.3. Compounds**

364 7-Deaza-2'-C-Methyladenosine (7-DMA) was purchased from Carbosynth (Berkshire,
365 UK) and dissolved in DMSO. β -D-N⁴-hydroxycytidine (EIDD-1931 or NHC) was

366 purchased from MedChemExpress (Monmouth Junction, NJ, USA) and dissolved in
367 DMSO.

368 **4.4. *Ae. aegypti* rearing**

369 *Ae. aegypti* Paea (Papeete, Tahiti, collected in 1994) were obtained via the Infavec2
370 consortium. For each rearing, eggs were hatched in dechlorinated tap water. After hatching,
371 groups of ± 400 larvae were transferred into trays containing 3 L of dechlorinated tap water
372 and fed every day with a yeast tablet (Gayelord Hauser, Saint-Genis-Laval, France) until
373 the pupae stage. Pupae were placed in small plastic containers inside cardboard cups for
374 their emergence. Adult mosquitoes were supplied with cotton balls soaked in a 10%
375 sucrose solution supplemented with 100 U/mL and 100 μ g/mL of PenStrep. Cardboard
376 cups containing adults were maintained at 28 ± 1 °C with a light/dark cycle of 16/8 h and
377 80% relative humidity.

378 **4.5. *Cx. pipiens* rearing**

379 *Cx. pipiens* biotype *pipiens* were kindly provided by Prof. Sander Koenraadt (Wageningen
380 University & Research, Wageningen, Netherlands) (15). For each rearing, eggs rafts were
381 hatched in trays containing 2 L of Milli-Q water (Synergy® UV, Merck, Germany). Larvae
382 were fed continuously until the pupae stage with TetraMin® baby fish food (Tetra,
383 Spectrum Brands, Germany). Pupae were collected as described in 4.4. Cardboard cups
384 containing adults were maintained at 25 ± 1 °C with a light/dark cycle of 16/8 h and 70%
385 relative humidity.

386 **4.6. Mosquito midgut dissection and *ex vivo* culture**

387 Unfed, antibiotic-treated female mosquitoes (3 – 7 days old) were cold-anaesthetized and
388 surface sterilized by soaking in 70% ethanol for 20 s followed by soaking in Dulbecco's

389 phosphate-buffered saline (PBS) for 20 s. Next, mosquitoes were dissected in PBS on a
390 petri dish using the stereomicroscope (VisiScope®, VWR). In brief, the midgut was
391 exposed by carefully pulling the second to last segment of the mosquito abdomen. The gut
392 of the mosquito was excised in its entirety (foregut, anterior and posterior midgut, and
393 hindgut) to keep the tissue of interest (midgut) from degradation, and tracheal tubes were
394 removed as much as possible without damaging the tissue. Following dissection, mosquito
395 guts were washed twice in midgut medium, consisting of: Leibovitz's L-15 medium
396 supplemented with 2% FBS, 100 U/mL of PenStrep, 50 µg/mL of kanamycin, and 0.25
397 µg/mL of amphotericin B (Sigma Aldrich, USA), and finally placed in a 96-well tissue
398 culture plate with a clear bottom (PerkinElmer®, USA) containing midgut medium. One
399 mosquito gut was placed in each testing well. Plates were maintained at 28°C, without CO₂.
400 Of note, handling and optimization of the *ex vivo* mosquito midguts culture are extensively
401 described in Supplementary Appendix 1.

402 **4.7. Videography of gut peristalsis and analysis**

403 *Ex vivo* cultured mosquito midguts were recorded with the Leica DMI8 microscope to
404 quantify the peristalsis observed *ex vivo*. A drop of carboxymethyl cellulose (CMC) 0.8%
405 diluted in Leibovitz's L-15 medium was deposited on a microscope glass slide and one
406 mosquito midgut was soaked in the drop. Forceps were used to gently arrange the hindgut
407 in a position suitable for analysis (horizontally positioned on the slide). Each hindgut was
408 recorded for a total of 1 minute 7 s (1 frame/0.3 s) and further discarded. This procedure
409 was repeated at several time points during incubation starting from day 0 (right after
410 dissection) to assess the contractility, and hence viability, of the *ex vivo* cultured midguts
411 over time.

412 For the analysis of the gut peristalsis, frames generated for each midgut were processed in
413 IgorPro (Wavemetrics, USA) using a custom script described for the measurement of
414 zebrafish gut motility (19). This script quantifies the changes in pixel intensity in a
415 designated area of analysis as it sequentially goes through all frames generated during the
416 video recording. As output, the software indicates individual peristaltic periodicity
417 (seconds in between contractions) per point of evaluation and overall averages of peristaltic
418 periodicity for each midgut analyzed. These outputs were annotated manually to check for
419 artifacts that could be generated by debris or tracheal tubes.

420 Serotonin hydrochloride (Sigma Aldrich, USA) and atropine sulfate salt monohydrate
421 (Sigma Aldrich, USA) were used to test the responsiveness of the mosquito midgut tissue.
422 Midguts were incubated with either serotonin (20 μ M), atropine (20 μ M), or midgut
423 medium for 1 hour, after which they were video recorded for further analysis.

424 **4.8. PrestoBlueTM (PB) assay**

425 The metabolic activity of the dissected midguts was further assessed using a resazurin salt-
426 based cell viability reagent PrestoBlueTM (Invitrogen, USA). For these experiments, three
427 dissected midguts were placed per well in a 96-well tissue culture plate. Each trio of
428 midguts was considered a biological replicate. Using more than one midgut per well was
429 necessary to ensure both a robust readout and a relatively rapid change of color in the
430 testing medium. Midguts fixated in 4% paraformaldehyde (PFA, Sigma Aldrich, USA)
431 were included in the assay as negative controls (NC). No organ (NO) controls were
432 composed of wells where one tip of the forceps was dipped in during dissection.

433 The live midguts and the corresponding controls were tested with PB every day for a period
434 of 10 days. In brief, the medium was carefully removed from the wells and replaced with

435 100 μ L of a 1:10 dilution of PB prepared in midgut medium (without phenol red). Next,
436 the midguts and their controls were incubated at 28°C without CO₂ for 7 hours. Once the
437 incubation time was completed, the PB-containing medium was transferred to a 96-well
438 tissue culture plate and fluorescence intensity was measured at wavelengths 560 nm
439 excitation and 590 nm emission with a Spark® Multimode Microplate Reader (Tecan
440 Trading AG, Switzerland). New midgut medium was added to wells containing midguts
441 and they were returned to the incubator (28°C, no CO₂) until the next time point
442 assessment. Two independent assays were carried out with at least 3 biological replicates
443 per time point.

444 **4.9. Infection and antiviral assays in *ex vivo* mosquito midguts**

445 Mosquito guts were dissected as described in 4.6 and placed into a 96-well tissue culture
446 plate filled with midgut medium. After removing the medium, a total of 1×10^4 PFU/mL
447 of the virus was added to each well. Of note, inocula of 1×10^5 PFU/mL and 1×10^6
448 PFU/mL were used for DENV-2 and USUV, respectively. Negative controls consisted of
449 dissected guts fixated in 4% PFA for 30 minutes. Midguts were incubated for two hours at
450 28°C, without CO₂. The virus inoculum was carefully removed, and midguts were washed
451 two times with midgut medium. Fresh medium was finally added to all wells for incubation
452 at 28°C, without CO₂.
453 To test the activity of antiviral drugs in the *ex vivo* mosquito midguts, compound dilutions
454 were prepared in midgut medium and added to the wells containing midguts to be treated,
455 after which they were infected with $1 \times 10^{4-5}$ PFU/mL of virus. Following two hours of
456 incubation at 28°C, the midguts were washed twice before adding midgut

457 medium alone (for virus control midguts) or containing compound (for treated midguts).

458 Midguts were returned to the incubator at 28°C, without CO₂.

459 Incubation time for alphavirus-infected midguts was 3 days, while flavivirus-infected
460 midguts were maintained for 7 days. Midguts were collected at several time points after
461 infection for further analysis.

462 **4.10. Determination of viral RNA levels and detection of infectious virus
463 replication**

464 Collected midguts were homogenized individually in 250 µL of PBS using bead disruption
465 (2.8 mm beads, Precellys). The midgut homogenate was filtered using 0.8 µm MINI
466 column filters (Sartorius, Germany) to remove debris, bacteria, and fungi. The filtered
467 homogenate was used for further viral RNA isolation and qRT-PCR to determine viral
468 RNA levels, and plaque assay to assess infectious virus particles.

469 Viral RNA isolation was performed with the NucleoSpin RNA Virus kit (Macherey-Nagel,
470 Germany) following the manufacturer's protocol. The sequences of primers and probes
471 used for each virus are compiled in Table 1. One-Step, quantitative RT-PCR was performed
472 for CHIKV and ZIKV in a total volume of 25 µL, consisting of 13.94 µL of RNase free
473 water (Promega, USA), 6.25 µL of master mix (Eurogentec, Belgium), 0.375 µL of each
474 forward and reverse primer (to a final concentration for each primer: 150 nM [CHIKV and
475 USUV]; 900 nM [ZIKV]), 1 µL of probe (to a final concentration of 400 nM [CHIKV and
476 USUV]; 200 nM [ZIKV]), 0.0625 µL of reverse transcriptase (Eurogentec, Belgium), and
477 3 µL of RNA sample. For RRV, the reaction mixture was prepared in a total volume of 20
478 µL, containing 5.2 µL of RNase free water, 10 µL of SYBR Green master mix (BioRad,
479 USA), 1 µL of each forward and reverse primer (to a final concentration of 125 nM for

480 each primer), 0.3 μ L of reverse transcriptase (BioRad, USA), and 4 μ L of RNA sample.

481 For DENV-2, the reaction mixture was prepared to a final volume of 20 μ L, consisting of

482 3 μ L of RNase free water, 10 μ L of SYBR Green master mix (BioRad, USA), 0.3 μ L of

483 each forward and reverse primer (to a final concentration of 900 nM for each primer), 0.4

484 μ L of reverse transcriptase (BioRad, USA) and 6 μ L of RNA sample.

485 The qRT-PCR assays were performed using the QuantStudioTM 5 Real-Time PCR System

486 (ThermoFisher Scientific, USA) with the following cycling conditions for CHIKV, ZIKV,

487 and USUV: 30 min at 48°C, 10 min at 95°C, followed by 40 cycles of 15 s at 95°C, and 1

488 min at 60°C (55 °C for USUV). Cycling conditions for RRV were as follows: 10 min at

489 50°C, 3 min at 95°C, followed by 40 cycles of 15 s at 95°C, and 30 s at 60°C. For DENV-

490 2, the following cycling conditions were used: 30 min at 48°C, 3 min at 95°C, followed by

491 40 cycles of 15 s at 95°C, and 1 min at 62°C.

492 For absolute quantification, standard curves were generated each run using 10-fold serial

493 dilutions of: cDNA of CHIKV-nsP1, viral RNA isolated from RRV and DENV-2 Bangkok

494 virus stocks or synthesized gBlocksTM gene fragments (Integrated DNA Technologies,

495 USA) for ZIKV and USUV.

496 Plaque assays were performed to quantify infectious virus particles from infected midgut

497 samples. Mammalian cells (Vero cells for RRV and CHIKV; BHK cells for ZIKV, DENV-

498 2, and USUV) were seeded at a density of 2.5×10^5 cells per well in 24-well tissue culture

499 plates. One-day confluent monolayers of cells were inoculated with serial 10-fold dilutions

500 of midgut samples, dilutions were prepared in 2% assay medium, and the inoculum was

501 allowed to infect the cells for 2 hours at 37°C, with 5% CO₂. A negative control consisting

502 of 2% assay medium used for preparing the dilutions was included in all plaque assays.

503 The inoculum was removed and replaced with CMC 0.8% overlay diluted in RPMI
504 medium (supplemented with 1% HEPES, 1% sodium bicarbonate, 1% L-glutamine, 1%
505 PenStrep and 2% FBS). Infected cells were incubated for 3 (*Alphaviruses*) or 7
506 (*Flaviviruses*) days. After incubation, 1 mL of 4% PFA was added to each well on top of
507 the overlay medium and allowed for fixation for 1 hour, after which the overlay mixture
508 was discarded, and the wells were carefully washed with water. The culture plates were
509 allowed to air dry, and then the wells were stained with 1% crystal violet solution (Sigma
510 Aldrich, USA). Plaques were counted and titer of each sample was calculated.

511 **4.11. Immunofluorescence staining of CHIKV viral proteins and imaging**

512 Mosquito guts were dissected and infected with CHIKV as described in 4.6 and 4.9,
513 respectively. At day 3 p.i., midguts were fixated with 4% PFA for one hour at room
514 temperature. Fixated midguts were transferred to a μ -slide 18-well (Ibidi GmbH, Germany)
515 plate and rinsed with PBS five times, for 5 minutes each time. Midguts were then saturated
516 with PBS-T (0.1% Triton X-100 [Sigma Aldrich, USA] and 1% bovine serum albumin
517 [BSA, Sigma Aldrich, USA] in PBS 1X) for 2 hours at room temperature. Primary antibody
518 anti-E2 protein [Chk265] (Absolute Antibody, Wilton, UK) was diluted in PBS-T (1:500)
519 and added to the fixed midguts for overnight incubation at 4°C. After incubation, midguts
520 were rinsed five times with PBS-T, for 5 to 10 minutes each rinse, followed by incubation
521 with Alexa Fluor 594 donkey anti-rabbit secondary antibody (ThermoFisher Scientific,
522 USA, A-21207, diluted in PBS-T, 1:500) for 2 hours at room temperature. Midguts were
523 rinsed five times with PBS-T, for 5 to 10 minutes each rinse. Phalloidin (diluted in 1%
524 BSA in PBS 1X solution, final concentration 100 nm) incubation followed for 10 minutes,
525 after which DAPI (diluted in PBS 1X, final concentration 100 nm) was added on top of the

526 phalloidin solution and allowed to incubate for 10 minutes. Finally, the phalloidin/DAPI
527 solution was removed, and midguts were rinsed with PBS. Midgut samples were mounted
528 in microscope slides using Glycergel® mounting medium (Agilent, USA) and allowed to
529 dry before imaging with a Leica DMi8 microscope (Leica Microsystems, Germany) and
530 the Andor Dragonfly Confocal Spinning Disk microscope (Oxford Instruments, UK). All
531 incubation steps were carried out on rotation, at 300 rpm.

532 **4.12. Infection of *ex vivo* cultured midguts with DENV-2 expressing
533 mCherry and imaging**

534 Mosquito guts were dissected and infected with a DENV-2 expressing mCherry (16) as
535 described in 4.6 and 4.9, respectively, using an inoculum of 1×10^5 PFU/mL. Negative
536 controls and mock-infected midguts were included. Mock-infected midguts were incubated
537 in midgut medium instead of virus dilution. Midguts were monitored every day during the
538 incubation period and checked under the FLoid™ Cell Imaging Station (Life
539 Technologies) for any signal that indicated mCherry expression. At day 7 p.i., the mosquito
540 midguts were fixated with 4% PFA for one hour at room temperature. After fixation,
541 midguts followed the protocol described in 4.11, skipping the addition of any primary and
542 secondary antibodies.

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563 **Supplementary material**

564 **Appendix I.** Handling and optimization of *ex vivo* mosquito midguts.

565 **Figure S1.** *Ex vivo* cultured mosquito guts were not significantly affected by external
566 incubation with serotonin or atropine.

567 **Figure S2.** Replication of DENV-2 expressing mCherry in the *ex vivo* cultured midguts.

568 **Figure S3.** ZIKV RNA loads in the mosquito midguts when cultured in medium or when
569 using a foam substrate for support.

570 **Figure S4.** CHIKV replication kinetics in mosquito midguts cultured in medium and in
571 carboxymethyl cellulose (CMC).

572 **Movie 1.** Representative mosquito hindgut videography used for the gut motility
573 analysis. Peristalsis is observed as waves along the hindgut region.

574

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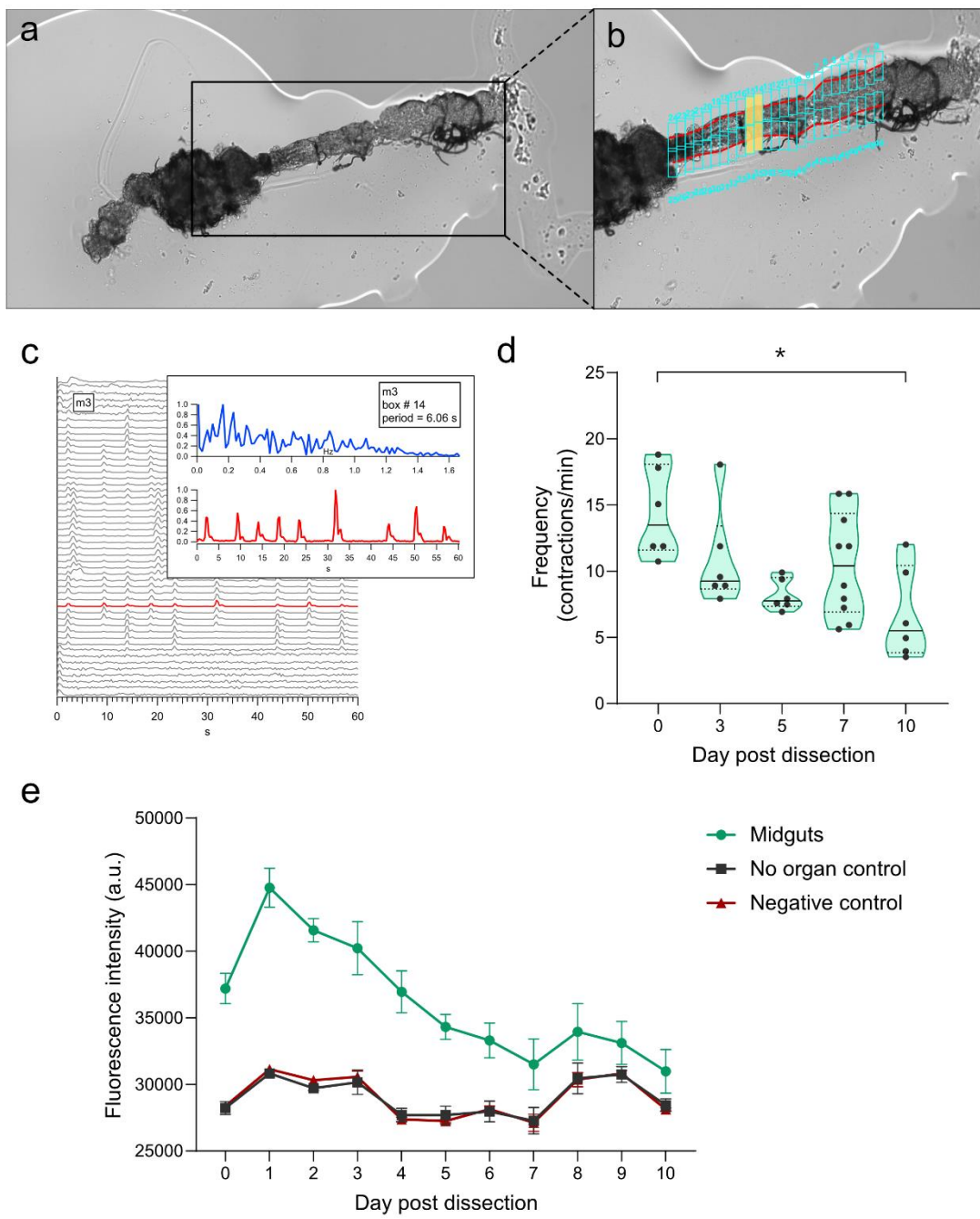
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700 **Figures and tables**

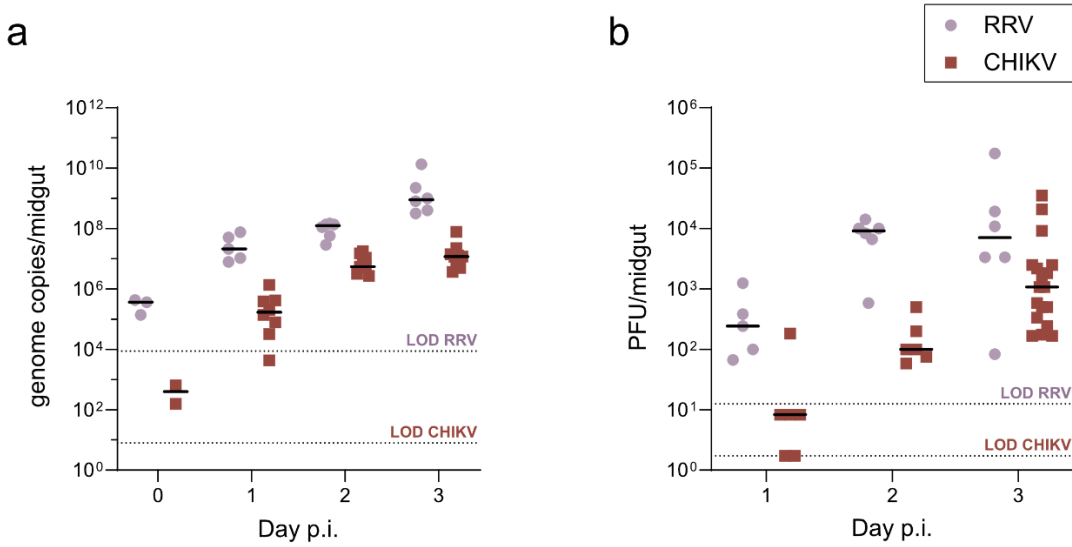


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Figure 1. *Ex vivo* cultured mosquito guts were viable for 7 days post dissection, based on gut peristalsis and a resazurin-based assay. **a**, Representative image of a mosquito gut placed in a carboxymethyl cellulose (CMC) drop for videography. **b**, Close-up of the hindgut which comprises the area of analysis (AOA) manually set and outlined by the red lines. Cyan boxes denoted the regions to be analyzed along the AOA. The number and size of the regions (cyan boxes) were adjusted to be 25 boxes on each side of the hindgut. **c**, Representative kymograph showing the output for one mosquito gut analysis. Each line in the kymograph corresponded to an individual region, which was further investigated in a plot. In the figure, the red highlighted line in the kymograph correlates to the presented plot displaying a representative analysis readout for this region. **d**, Contractions observed per minute in function of time post dissection in days. Each point represents one individual mosquito midgut. The black line represents the median value. The graph shows data from three independent experiments. Statistical significance was assessed with a Kruskal-Wallis test. Significantly different values are indicated by asterisks: *, P <0.05. **e**, Viability readouts exhibit metabolic activity for a period of 10 days for tested midguts, no organ and negative controls. Assays were performed with 2 or 3 biological replicates, each replicate composed of 3 midguts per well, along with its corresponding controls. Error bars represent the standard errors of the means per time point. The graph shows data from three independent assays.

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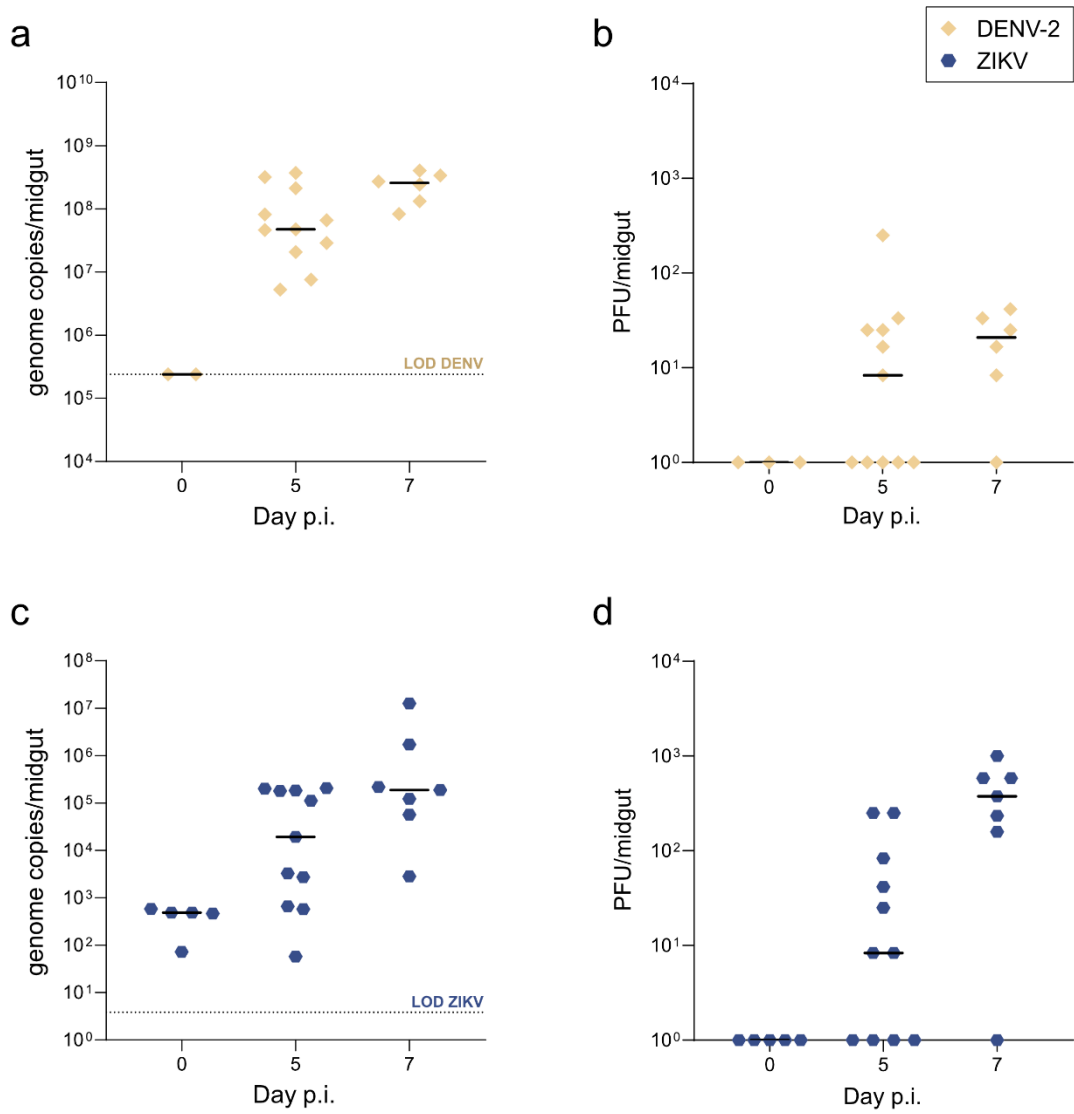
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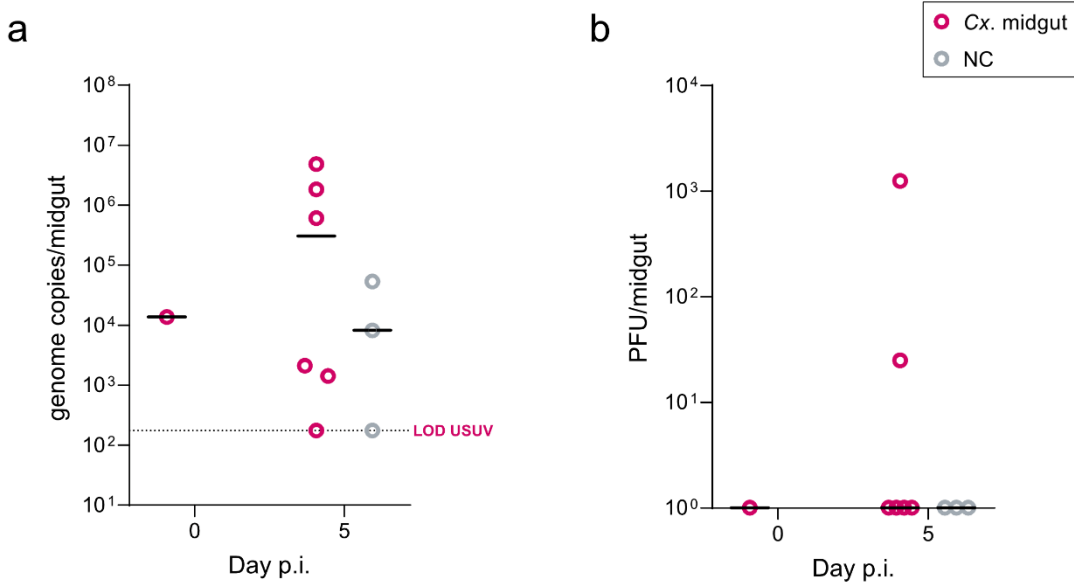
Figure 2. *Ex vivo* cultured midguts supported Ross River (RRV) and chikungunya virus (CHIKV) infection. a, Viral RNA levels in the mosquito midguts were quantified at 2 hours (day 0), 1, 2 and 3 days p.i. by means of qRT-PCR. b, Infectious virus loads in the mosquito midguts were quantified by means of plaque assay. Each dot represents an individual midgut organ. The black line represents the median value. Data correspond to at least two independent replication kinetics assays. LOD: Limit of detection of the corresponding assay.

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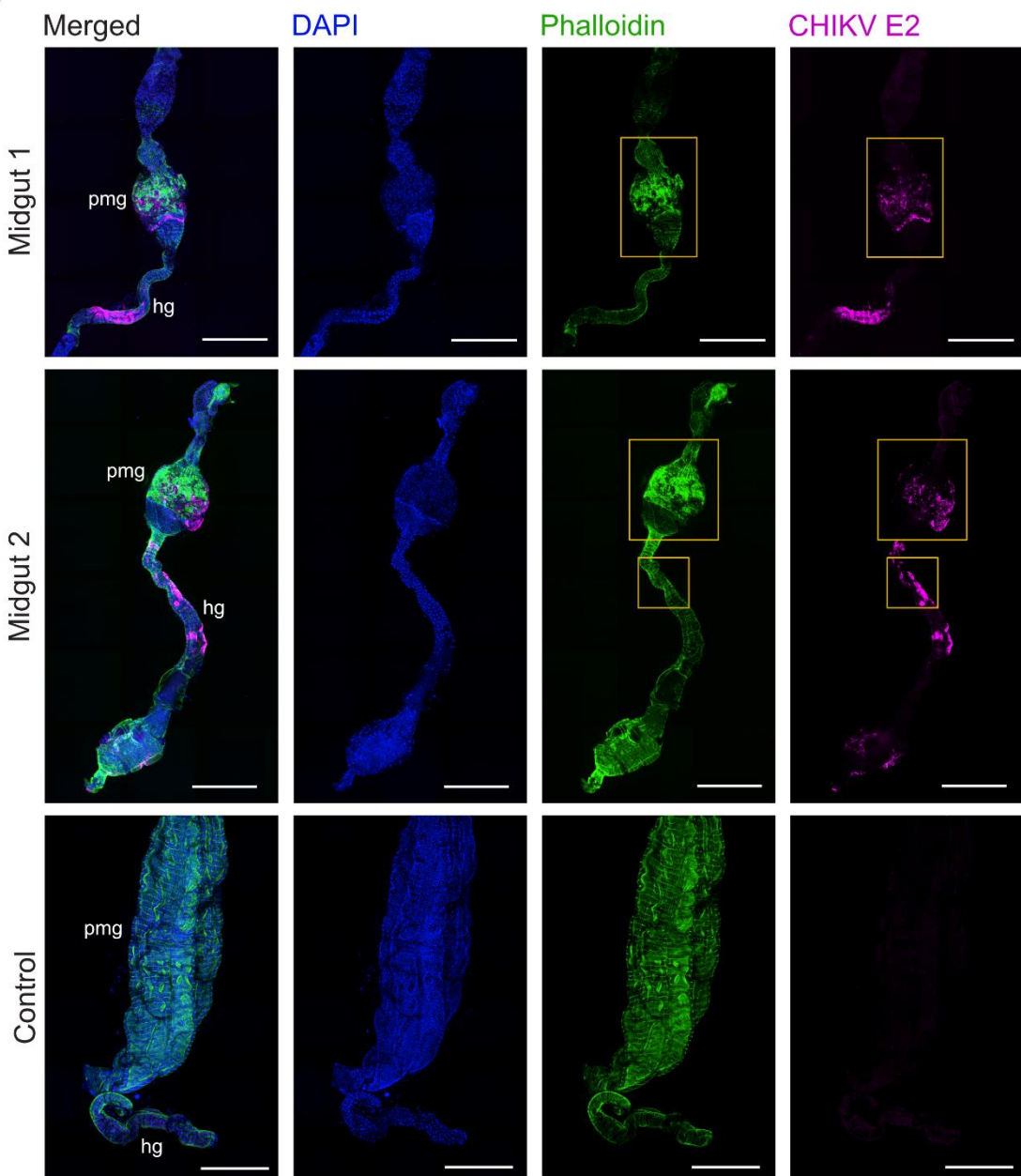
Figure 3. *Ex vivo* cultured midguts supported dengue virus serotype 2 (DENV-2) and Zika virus (ZIKV) infection. **a, c**, Viral RNA levels in the mosquito midguts were quantified at 2 hours, 5 and 7 days p.i. by means of qRT-PCR. **b, d**, Infectious virus loads in the mosquito midguts were quantified by means of plaque assay. Each symbol represents an individual midgut organ. The black line represents the median value. Data correspond to two independent replication kinetics assays. LOD: Limit of detection of the corresponding assay.



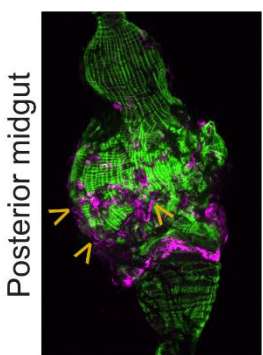
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Figure 4. *Ex vivo* cultured midguts were partially permissive to Usutu virus (USUV) infection. **a**, Viral RNA levels in the mosquito midguts were quantified at 2 hours and 5 days p.i. by means of qRT-PCR. **b**, Infectious virus loads in the mosquito midguts were quantified by means of plaque assay. Each symbol represents an individual midgut organ. The black line represents the median value. LOD: Limit of detection.

a



b



c

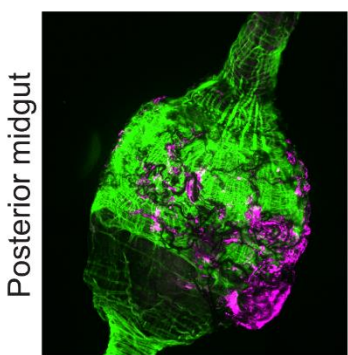


Figure 5. Detection of CHIKV protein synthesis in infected *ex vivo* midgut cultures. **a**, Magnification, 25X. Overlay of blue, green, and magenta filter imaging are shown for two infected midgut organs and one negative control midgut. Imaging of each individual channel as follows: in blue, DAPI-stained cell nuclei; in green, actin filaments stained with phalloidin; and in magenta, E2 viral protein. No E2 expression was detected in the negative control midguts. The scale bar in panels, represented by the white line, corresponds to 400 μ M. pmg: posterior midgut. hg: hindgut. **b**, Close-up panel showing infection foci in the posterior midgut region corresponding to the yellow squares indicated in **a** for “Midgut 1”. Arrow heads indicate E2 expression localized in tracheal tubes of the midgut. **c**, Close-up panel showing infection foci in the posterior midgut and hindgut region corresponding to the yellow squares indicated in **a** for “Midgut 2”.

709

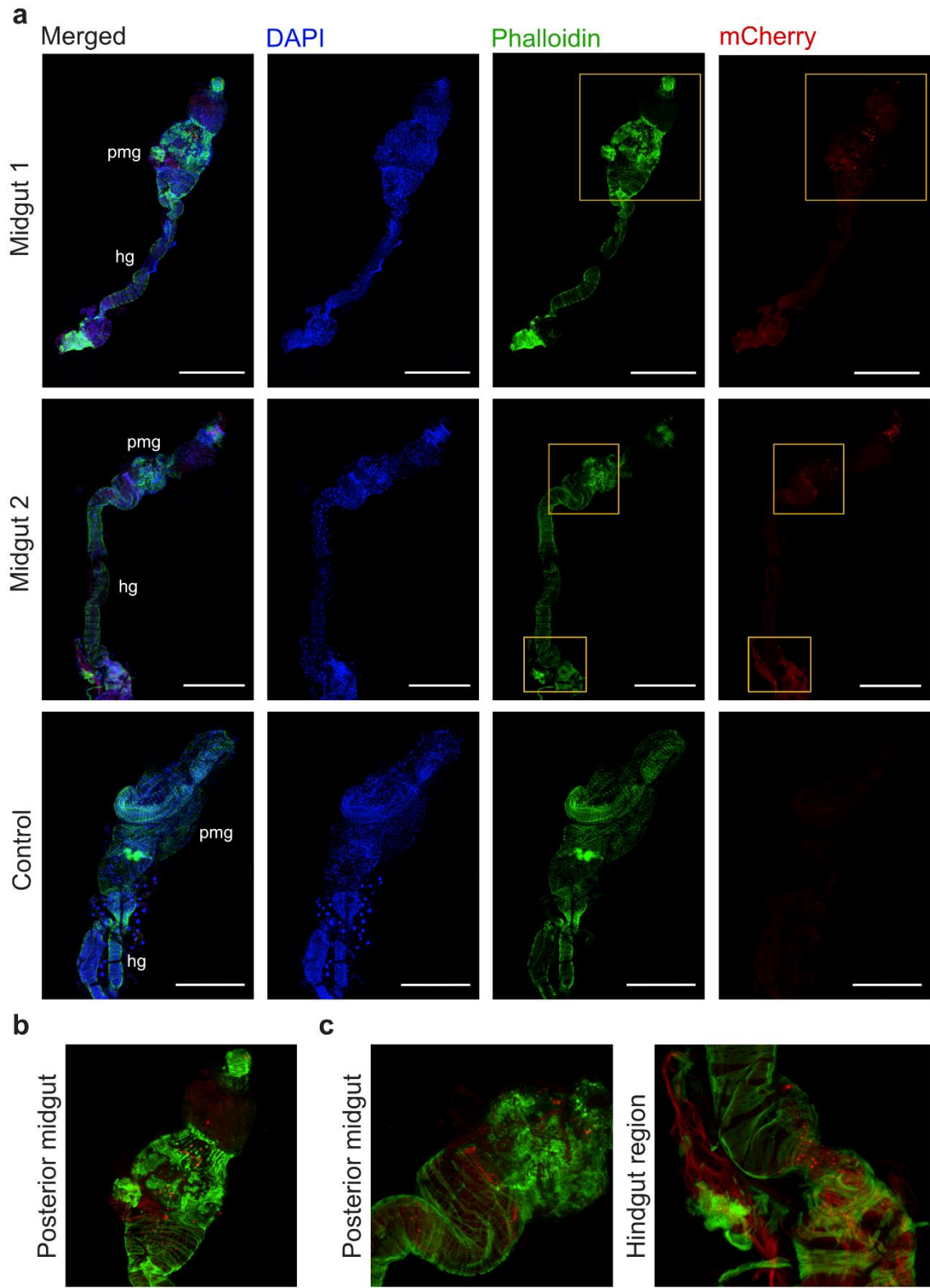
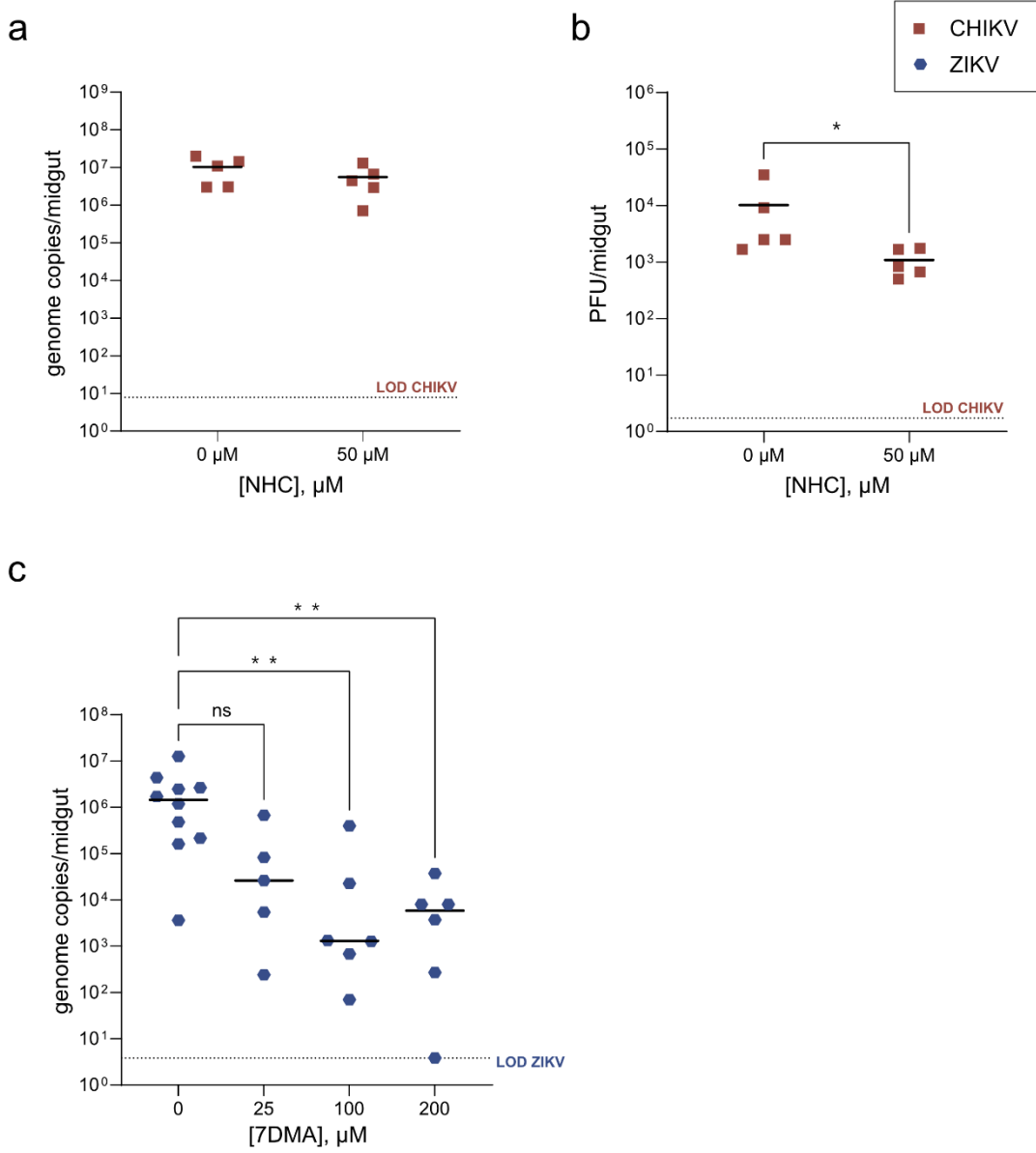


Figure 6. Replication of DENV-2 expressing mCherry in the *ex vivo* cultured midguts. **a**, Magnification, 25X. Confocal microscopy displays the DENV-2 infection in the *ex vivo* midguts at day 7 p.i., as seen by the mCherry (red) signal. Overlay of blue, green, and red filter imaging are shown for two midgut organs and one negative control midgut. Imaging of each individual channel as follows: in blue, DAPI-stained cell nuclei; in green, actin filaments stained with phalloidin; and in red, mCherry signal. No mCherry expression was detected in fixated or mock-infected midguts. The scale bar in panels, represented by the white line, corresponds to 400 μ M. pmg: posterior midgut. hg: hindgut. **b**, Close-up panel showing several infection foci in the posterior midgut region corresponding to the yellow squares indicated in **a** for “Midgut 1”. **c**, Close-up panel showing infection foci in the posterior midgut and hindgut region corresponding to the yellow squares indicated in **a** for “Midgut 2”.

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Figure 7. Antiviral activity of NHC and 7DMA in the *ex vivo* cultured mosquito midguts. **a**, CHIKV RNA loads in the mosquito midguts were quantified at day 3 p.i. by qRT-PCR. **b**, CHIKV infectious virus loads were quantified at day 3 p.i. by plaque assay. Statistical significance was assessed with a Mann-Whitney test. Significantly different values are indicated by asterisks: *, P<0.05. **c**, ZIKV RNA levels in the mosquito midguts were quantified at day 7 p.i. by qRT-PCR. Statistical significance was assessed with a Kruskal-Wallis test. Significantly different values are indicated by asterisks: **, P<0.05. ns: not significant. Each dot represents an individual midgut. The black line represents the mean value. LOD: Limit of detection of the corresponding assay.

714

Table 1. Sequences of primers and probes for qRT-PCR used in this study

Virus [target protein]	Forward primer (5' → 3')	Reverse primer (5' → 3')	Probe (5' → 3')	Reference
Ross River virus [E2]	TACAAGCACGACCCATTGCCG	GATAGTCCTGCCGCCTGCTGT	N/A	(36)
Chikungunya virus [nsP1]	CCGACTCAACCATCCTGGAT	GGCAGACGCAGTGGTACTTCCT	FAM-TCCGACATCATCCTCCTGCTGGC-TAMRA	(37)
Zika virus [E]	CCGCTGCCAACACAAG	CCACTAACGTTCTTTGCAGACAT	FAM-AGCCTACCT-ZEN-TGACAAGCAATCAGACACTCAA-IABkFQ	(38)
Dengue virus serotype 2 [NS3]	CAGATGGAGCTGGAGTTAC	TTTGACGTCCGCCATGAA	N/A	<i>In-house</i>
Usutu virus [NS5]	AAAAATGTACGCGGATGACACA	TTTGGCCTCGTTGTCAAGATC	FAM-TGGGACACCCGGATACCAGAG-TAMRA	(39)

