1 Experimental adaptation of dengue virus 1 to Aedes albopictus

2 mosquitoes by *in vivo* selection

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22 Abstract

23 In most of the world, Dengue virus (DENV) is mainly transmitted by the mosquito Aedes 24 aegypti while in Europe, Aedes albopictus is responsible for human DENV cases since 2010. 25 Identifying mutations that make DENV more competent for transmission by Ae. albopictus 26 will help to predict emergence of epidemic strains. Ten serial passages in vivo in Ae. 27 albopictus led to select DENV-1 strains with greater infectivity for this vector in vivo and in 28 cultured mosquito cells. These changes were mediated by multiple adaptive mutations in the 29 virus genome, including a mutation at position 10,418 in the DENV 3'UTR within an RNA 30 stem-loop structure involved in subgenomic flavivirus RNA (sfRNA) production. Using 31 reverse genetics, we showed that the 10,418 mutation alone does not confer a detectable 32 increase in transmission efficiency in vivo. These results reveal the complex adaptive 33 landscape of DENV transmission by mosquitoes and emphasize the role of epistasis in 34 shaping evolutionary trajectories of DENV variants.

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Keywords: Vector-borne diseases, Europe, Emergence, Invasive species, Dengue, *Ae*.
 albopictus, adaptive-evolution, sfRNA

38

40 Introduction

41 Vector-borne diseases represent almost one fourth of all emerging infectious diseases 42 worldwide (1). Among the emerging diseases, arboviruses occupy the top stair with several 43 million human cases reported annually (2). Dengue virus (DENV) belongs to the Flavivirus 44 genus of the *Flaviviridae* family and is by far the most important arboviral disease, with the 45 number of human dengue infection cases exceeding 300 million annually; 96 million are symptomatic dengue fever/hemorrhagic fever leading to an estimated 22,000 human deaths 46 (3). DENV is comprised of four antigenically distinct but genetically related serotypes 47 48 referred to as DENV1-4 (4). Clinical manifestations range from mild cases of dengue fever to 49 severe cases of dengue hemorrhagic fever and/or dengue shock syndrome. All four DENV 50 serotypes are now circulating in Asia, Africa and America (5). In past centuries, dengue was not an uncommon disease in Europe: the last record of a dengue outbreak in the 20th century 51 52 was in Athens, Greece, in 1927–1928 (6). This outbreak was unusual by the number of cases 53 (~1 million) and the importance of severe clinical symptoms (e.g. hemorrhagic 54 manifestations) leading to deaths (~1,000). After this Greek episode, dengue disappeared 55 from Europe (7) as the mosquito Aedes aegypti disappeared from Eastern Mediterranean after 56 1935 through improving sanitation and mosquito control measures (8). No local transmission 57 of DENV has been reported in Europe until 2010, when clusters of autochthonous cases were 58 reported in Southern France (9) and Croatia (10). In France, several transmission episodes 59 were successively reported: 2013-2015 (11-13), 2018-2019 (14, 15). The vector was Aedes 60 albopictus, first detected in Europe in 1979 in Albania (16), then in 1990 in Italy (17), and 61 today, established in more than 20 European countries (18). Unexpectedly, Ae. albopictus 62 from France was shown to be more competent to experimentally transmit DENV-1 strains 63 compared to its counterpart Ae. aegypti from the French West Indies (19). Contrary to Ae. 64 *aegypti*, the mosquito Ae. albopictus which is native to South-East Asia, has a broader range 65 of hosts (20). When Ae. aegypti is absent, Ae. albopictus can be responsible for DENV

epidemics, as shown for the outbreaks in the Seychelles islands (21), Japan (22), La Réunion
Island (23), and Hawaii (24). However, to date, *Ae. albopictus* is considered a minor vector of
DENV relative to *Ae. aegypti* (25).

69 DENV serotypes have caused recent epidemics by changing their host ranges to 70 increase infections in humans (26). As most arboviruses, DENV is capable of rapidly 71 adapting to changes in their environment (or novel hosts) due to the accumulation of one or 72 more specific mutations in the viral genome. For DENV, mutations in the 3' untranslated 73 region (UTR) have previously been linked to increased epidemiological fitness of the virus 74 via a mechanism involving increased expression of 3' UTR-derived subgenomic flavivirus 75 RNA (sfRNA) expression (27). SfRNA is a known determinant for mosquito transmission of 76 multiple flaviviruses like DENV, Zika virus and West Nile virus (28-32). Nucleotide 77 substitutions in the 3'UTR reducing or ablating sfRNA expression negatively impact viral 78 infection and transmission rates, suggesting that there is evolutionary pressure on 79 conservation of RNA structures that dictate sfRNA expression in mosquitoes (33, 34). These 80 studies indicate that subtle changes in the viral nucleotide composition can enhance the viral 81 epidemic potential. On the same line, CHIKV has acquired the ability to spread globally 82 owing to a single Ae. albopictus-adaptive mutation E1-226V (35). This mutation increased the 83 infectivity of CHIKV in Ae. albopictus (36, 37).

84 We hypothesize that DENV can be selected for enhanced transmission by European Ae. 85 *albopictus*, which would provide insight into future epidemic DENV strains that could pose a 86 threat to human health. We conducted an experimental evolution study to identify nucleotide 87 changes in the DENV genome by serially passaging DENV-1 isolates from Thailand (30A) 88 and France (1806) in an Ae. albopictus population from Nice, France. Ten total passages were 89 completed after which viral isolates were deep sequenced to identify newly acquired 90 mutations. Importantly, we investigated whether the adaptation to the mosquito vector 91 resulted in enhanced transmission potential or replication rate in mosquitoes. These results

92 exemplify the potential of virus-adaptation studies for the identification of DENV strains

93 likely to emerge.

94

95 Results

96 European *Ae. albopictus* are differentially susceptible to DENV-1.

97 Arboviral transmission requires competent mosquitoes. To test whether European populations 98 of Ae. albopictus can sustain local transmission of DENV-1, as reported in France (9) and 99 Croatia (10), Ae. albopictus mosquitoes from Alessandria and Genoa (Italy), Cornelia and 100 Martorell (Spain), Nice and Saint-Raphael (France) were experimentally infected with 101 DENV-1 1806 from France or with DENV-1 30A from Thailand. Only engorged females 102 were kept for analysis (samples size indicated in Fig. 1). When examining viral infection rate 103 (Fig. 1a) and dissemination efficiency (Fig. 1b) at 14 and 21 days post-infection (dpi), 104 percentages increased along with the dpi for the majority of virus/mosquito combinations. 105 Within a mosquito population, the viral strain did not play a major role in either infection rate or dissemination efficiency (Fisher's exact test: p > 0.05 after Bonferroni correction; Table 106 107 S1). In contrast, we observed significant differences between mosquito populations (Fisher's 108 exact test: p < 0.05; Table S2) meaning that the geographic origin of the mosquito population 109 is a critical factor that determines the outcome of viral infection and dissemination. Viral 110 loads in heads (indicative of a successful dissemination from the midgut) did not mostly differ among mosquitoes having disseminated the virus (Fig. 1c, Table S3) (Wilcoxon Rank-Sum 111 112 test: p > 0.05). No viral particles were detected in mosquito saliva (data not shown).

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114 Experimental adaptation of DENV-1 to Ae. albopictus.

To examine whether DENV-1 can adapt to *Ae. albopictus*, DENV-1 1806 and DENV-1 30A
were passaged 10 times in duplicate in *Ae. albopictus* mosquitoes from Nice, France (Fig. 2).
Viral titers of cell culture supernatants collected at each passage fluctuated slightly from

passages 1 to 10 (Fig. S1). Additionally, the viruses were passaged 10 times in duplicate in 118 119 Ae. albopictus C6/36 cells as a cell culture control. Full viral genomes were examined by 120 deep sequencing at each passage (1-10) for the two replicates (R1 and R2) of *in vivo* mosquito 121 infections and for passages 0 (parental strain), 1, 5 and 10 for the C6/36 cell culture control. 122 The two parental DENV-1 strains, 1806 and 30A, yielded a mean sequencing depth of 123 68,687X (1806) and 133,941X (30A), covering 99.97% and 100% of the reference genome at 124 >100X. With the exception of 30A (passage 4, R2 in mosquitoes; covering only 34.62% of 125 the reference genome at >100X), all passages had a mean coverage between 995X and 126 211,830X, paving between 100 and 99.25% of the reference genome at >100X (Fig. S2). 127 No major changes in single nucleotide variants (SNV) frequencies were detected when 128 DENV-1 isolate was serially passaged on C6/36 cells (Fig. 3a). Remarkably, we did not 129 detect a single mutation that reached consensus level (frequency >50%) in the C6/36 control 130 passages. In contrast, when DENV-1 1806 or 30A was passaged in Ae. albopictus 131 mosquitoes, consensus level variants were detected as soon as passages 2 (DENV-1 1806) and 132 5 (DENV-1 30A), and a total of 30 consensus level variants were detected at passage 10 (Fig. 133 3a, Table S4). In total, twenty consensus level SNVs were detected in DENV-1 30A 134 (positions 448, 694, 1611, 1768, 1959, 2002, 2200, 2716, 2977, 3442, 5822, 6658, 6728, 135 7267, 7952, 8149, 8485, 9504, 10208, 10258) and 10 in DENV-1 1806 (positions 1840, 2719, 3001, 3757, 4552, 4606, 5667, 7360, 9067, 10418). Out of the 30 consensus variants, 23 136 137 synonymous changes, 6 non-synonymous and one variant located in the 3'UTR were detected 138 (Fig. 3b). The variant located at position 10,418 in the 3'UTR was the only SNV shared 139 between the replicates R1 and R2 of DENV-1 1806. Its frequency increased over the 140 passages, reaching consensus level at passage 4 for replicate 1 and passage 8 for replicate 2 141 (Fig. 3a). The variant became almost fixed (frequency > 99%) at passage 5 for replicate 1 and 142 passage 10 in replicate 2. No SNV was common to the two replicates for DENV-1 30A. 143 These results indicate that DENV-1 accumulates mutations during passaging in Ae. albopictus

that likely facilitate virus replication in the mosquito or virus dissemination into the saliva tofacilitate transmission.

146

147 Ae. albopictus adapted DENV-1 1806 has an increased transmission rate in Ae. 148 albopictus.

To investigate whether the mosquito adapted DENV-1 1806 has increased transmission 149 potential as compared to the parental isolate, Ae. aegypti Pazar (Turkey) and Ae. albopictus 150 Nice (France) were provided with an infectious blood-meal containing 10⁷ FFU/mL DENV-1 151 152 1806 parental, or replicate of the mosquito passaged virus (R1 and R2). Viral infection rates 153 were high (>40%) and higher for the P10 viruses in Ae. albopictus at 21 dpi (Fig. 4a-b). Viral 154 dissemination was lower at early dpi but remained high at 21 dpi in both mosquito species 155 with a higher dissemination of P10 viruses compared to the parental virus (Fig. 4c-d). Transmission was surprisingly low in Ae. aegypti (Fig. 4E) compared to Ae. albopictus (Fig. 156 157 4f) suggesting a stronger effect of salivary glands as a barrier to virus release in saliva of Ae. 158 aegypti. At 21 dpi, the two P10 viruses had higher infection rates (Fig. 4b), higher 159 dissemination rates (Fig. 4d) and higher transmission rates (~2.5 fold) in Ae. albopictus than their parental strain (Fisher's exact test: $p < 10^{-4}$; IR, p = 0.0001; DE, p = 0.0001; TE, p =160 161 (0.022). Collectively, these results indicate that the accumulation of adaptive mutations during 162 passaging in Ae. albopictus for the P10_R1 and P10_R2 viruses is beneficial for virus 163 infection, dissemination and thus transmission by Ae. albopictus mosquitoes.

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Ae. albopictus adapted DENV-1 1806 has a replicative advantage in RNAi-competent and -deficient mosquito cells.

To determine whether the adaptive mutations in DENV-1 1806 after serial passaging in *Ae*. *albopictus* mosquitoes were causing a replicative advantage, we examined the replication
kinetics of DENV-1 1806 parental, R1 and R2 in *Ae. albopictus* C6/36 (RNAi-deficient) and

170 U4.4 (RNAi-competent) cells (Fig. 5a-b) compared to kinetics in HFF cells (Fig. 5c). In 171 C6/36 cells, the parental virus reached slightly lower titers at 6, 24 and 48 hours post-172 infection (hpi) as compared to the mosquito passaged R1/R2 viruses (Fig. 5a). The mosquito 173 adapted R1/R2 viruses presented a significant increase in viral titer at 24 hpi (R1 (mean \pm 174 SD): $Log_{10} 4.42 \pm 0.10$; R2: $Log_{10} 4.68 \pm 0.14$) compared to the parental strain ($Log_{10} 3.31 \pm 0.14$) 175 0.15) (χ^2 test: p = 0.027). In U4.4 cells, the same trend was observed (Fig. 5b) with a lower 176 titer at 24 hpi for the parental strain ($Log_{10} 2.84 \pm 0.06$) as opposed to the two P10 viruses 177 (R1: $Log_{10} 4.19 \pm 0.29$; R2: $Log_{10} 4.47 \pm 0.03$). In human cells (Fig. 5c), viral titers remained 178 between 2 and 3 Log_{10} from 0 to 72 hpi. These results indicate that the adaptive mutations 179 after serial passaging of DENV-1 1806 in Ae. albopictus mosquitoes increase the replication 180 rate in mosquito cells and a disadvantage in human cells.

181

182 Vizualisation of substitutions in 3'UTR of DENV-1 1806 on RNA stem-loop structures.

183 The highly structured flavivirus 3'UTR is important for virus replication, genome translation 184 and production of non-coding sfRNA (38). sfRNA is formed as a result of incomplete 185 degradation of the viral genomic RNA by the 5'-3' exoribonuclease XRN1, which stalls on 186 stem loop (SL) and dumbbell (DB) RNA structures in the 3'UTR (38, 39). It has been shown 187 that passaging DENV on mosquito cells can result in high mutation rates in the 3'UTR and 188 might alter the abundance of sfRNA during infection (33, 40). The largest sfRNA species, 189 sfRNA1, determines pathogenicity (41), inhibits host innate immunity (27, 42) and is 190 essential for efficient transmission of flaviviruses by mosquitoes (28-32, 43, 44). We 191 therefore investigated if the consensus level mutation 10,418 that occurred in the 3'UTR after 192 passaging could lead to changes in the 3'UTR secondary RNA structures and subsequent 193 sfRNA formation. Mutations with an SNV frequency ≥ 0.05 only occurred in the SL-II and 194 3'SL structures (Fig. 6a; red nucleotides). When examining mutations in the 3'SL, the same 195 SNVs were found in the two parental strains and the passages 10 indicating that those

196 mutations were already present in the initial viral populations and were not selected 197 consequently to serial passages in mosquitoes (Fig. 6b). We observed that the mosquito 198 passaged DENV-1 1806 presented a U \rightarrow C substitution at position 10,418 on the top of SL-II, 199 which was not observed for DENV-1 30A. SL-II is the XRN1 stalling structure required for 200 sfRNA2 formation, which requires the presence of a RNA pseudoknot interaction and a 201 complex tertiary folding (45) (Fig. 6b). Pseudoknot formation and other known tertiary RNA 202 interactions are not expected to be directly disrupted due to the 10,4018 sequence change 203 (Fig. 6b). However, the U \rightarrow C mutation may indirectly affect the 3D folding of the stem loop. 204 Passages on C6/36 cells also gave rise to lower-frequency mutations in SL-II (frequency <205 0.2), but none of them reached consensus level.

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Increased transmission potential of *Ae. albopictus*-adapted DENV-1 is not associated with significantly increased sfRNA production.

209 As sfRNAs are generated due to stalling of XRN1 on RNA secondary structures in the viral 210 3'UTR (41), sequence changes in the 3'UTR, in particular those that occur in RNA structures 211 involved in XRN1 stalling, may affect the length and expression level of sfRNAs. To investigate whether the production of sfRNA is affected by the mutation 10,418 in DENV-1 212 213 1806 R1/R2, a Northern blot analysis was performed using a 3'UTR specific probe on total 214 RNA extracted from U4.4 cells infected with DENV-1 1806 parental, R1 or R2 (Fig. 7). Viral 215 gRNA and abundant sfRNA1 were produced by both the parental and R1/R2 viruses (Fig. 7a). 216 The quantity of sfRNA1 was visually similar across all samples on the gel, although ImageJ 217 quantification of the band intensities revealed that the ratio of sfRNA/gRNA was ~1.5 fold 218 higher in R1 and R2 samples as compared to the parental samples (Fig. 7b). Minimal amounts 219 of smaller sfRNA species (i.e. sfRNA2, 3, 4) were observed, indicating that DENV-1 220 predominantly produces sfRNA1 during infection of mosquito cells. These results show that

the increased transmission potential of the *Ae. albopictus* adapted DENV-1 1806 is unlikely to

222 be caused by differences in sfRNA production.

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224 The 10,418 mutation alone does not significantly enhance DENV-1 transmission by Ae.

225 *albopictus*

226 To test whether the U \rightarrow C substitution at position 10,418 alone would recapitulate the 227 observed phenotype in Ae. albopictus, three reverse genetic constructs (Parental construct, 228 P10 construct 1 and P10 construct 2) were produced using the ISA method and then 229 sequenced. As expected, the two P10 constructs presented the 10,418 mutation at a frequency 230 close to 100% (Table S5). The genetic constructs also displayed other SNVs at frequencies 231 higher than 5% but none of them reached consensus level with the exception of one SNV with 232 52.5% frequency in P10 construct 1 (5173/nsp3) and two SNVs close to fixation in P10 233 construct 2 (7321/nsp4b and 9571/nsp5).

234 Twenty-one days after an infectious blood meal containing the reverse genetic 235 constructs, Ae. albopictus Nice mosquitoes were examined for transmission by analyzing viral 236 particles in their saliva. When estimating the transmission efficiency, no significant 237 differences were detected when comparing the five viral strains, the reverse genetic constructs 238 in reference to the template (Parental, P10) and the two P10 constructs (1 and 2) (Fisher's 239 exact test: p > 0.05) (Fig. 8a). Similarly, when examining the number of viral particles in 240 individual mosquito saliva, no statistical significance was found whatever the comparison 241 (Wilcoxon Rank-Sum test: p > 0.05) (Fig. 8b).

To confirm the profile of P10 construct 1, we performed a replicate using the same experimental design. TE (Fig. 8c) and the viral load in saliva (Fig. 8d) were determined. We found that the replicate 2 shares the same profile than the replicate 1 (Fig. 8a, 8b). Altogether, these results indicate that the mutation 10,418 alone does not enhance transmission of DENV-1 in *Ae. albopictus*.

247

248 Discussion

249 In this study, we developed an experimental evolution approach to enhance the transmission 250 of DENV-1 by the vector Ae. albopictus. A number of conserved nucleotide variants were 251 observed in all mosquito-passaged DENV-1 viruses, but not in the viruses passaged in 252 mosquito cells, indicating that DENV-1 virus adapts specifically to cope with the adaptive 253 pressure in the mosquito. A nucleotide change at position 10,418 in the 3'UTR was observed 254 in both replicates of the Ae. albopictus adapted DENV-1 1806. Independent fixation of the 255 mutation in two replicates with a sharp rise in frequency is consistent with adaptive evolution. 256 Importantly, the Ae. albopictus adapted isolates displayed higher DENV-1 infection, 257 dissemination and transmission efficiencies in Ae. albopictus. Our results show that we have successfully adapted DENV-1 to Ae. albopictus, through selection of adaptive mutations 258 259 including the 10,418 mutation in the 3'UTR of the viral genome by sequential passaging in 260 vivo.

261 Ae. albopictus usually acts as a secondary vector of DENV (20), but in the absence of 262 Ae. aegypti, it can act as the main vector in some regions including Europe (9-13). First 263 detected in Albania in 1979 (16), Ae. albopictus is now present in more than 20 European 264 countries (18). We showed that Ae. albopictus from France, Italy and Spain were susceptible 265 to infection by DENV-1 (Fig. 1), indicating that Ae. albopictus is indeed a vector species for 266 DENV in Europe. The main sources of introductions in Europe were mosquitoes from Italy, 267 which were previously imported from North America (46). Recurrent introduction events 268 have contributed to increase the genetic diversity of European Ae. albopictus populations 269 (47), an important factor shaping vector competence (25).

270 Here, we experimentally selected DENV-1 isolates for enhanced transmission by *Ae*.
271 *albopictus*. Our experimental procedure was designed to accelerate the selection process of
272 DENV-1 by serial passages in *Ae. albopictus* mosquitoes without alternation in the

273 mammalian host. After 10 passages in Ae. albopictus collected in Nice, France, we 274 successfully adapted DENV-1 1806 and DENV-1 30A to Ae. albopictus through the 275 accumulation of adaptive mutations across the genome, although only a single mutation was 276 fixed in both replicates for the 1806 isolate. Importantly, for DENV-1 1806, these adaptive 277 mutations increased the infection, dissemination and transmission rates of DENV-1 by Ae. 278 albopictus (Fig. 4). Furthermore, growth kinetics of the DENV-1 1806 viruses were increased 279 in both RNAi-competent U4.4 and RNAi-deficient C6/36 cells, indicating that the mutations 280 cause an increase in viral replicative fitness in cell cultures regardless of a functional RNAi 281 machinery (Fig. 5). These mosquito-selected viral variants were less adapted to replicate on 282 mammalian cells (48). In a similar approach, Stapleford et al. (2014) succeeded in monitoring 283 the selection of epidemic variants of CHIKV adapted to Ae. albopictus consolidating the idea 284 that in vivo approaches can contribute in predicting new variants able to emerge and displace 285 currently circulating viral strains (49).

286 We identified a mutation that was fixed in both replicates of the mosquito passaged 287 DENV-1806 isolate, suggesting that this residue is involved in the adaptive-evolution that 288 results in the increased transmission and replication potential of the virus. This shared 289 substitution was located at position 10,418 in the highly structured 3'UTR of the DENV-1 290 genome. Specifically, the mutation was present in the SL-II RNA structure (Fig. 6); which is 291 required for the production of sfRNA2 (41). XRN1 stalls at SL and dumbbell (DB) RNA 292 structures within the 3'UTR, which results in accumulating sfRNAs of different sizes (41, 293 50). The stalling of XRN1 occurs due to steric hindrance caused by interactions of 294 pseudoknots (PK) and other tertiary RNA structures (45, 51). Prediction of RNA structures 295 involved in XRN1 stalling (the so-called xrRNAs) with Mfold has proven to be an useful 296 starting point but undeniably has limitations, e.g. pseudoknots cannot be predicted and 3D 297 RNA folding is not taken into account. Although Mfold predictions and visual pseudoknot 298 mapping have helped to elucidate mechanisms of XRN1 stalling (41, 45, 50), the exact

299 structural basis for XRN1 stalling, the involvement of a unique three-way junction and 300 internal tertiary interactions were only revealed by determining the crystal structure of several 301 xrRNAs (52, 53). In mammalian cells, sfRNA is essential for inducing pathogenicity (41), 302 and acts as an antagonist of innate immune responses (27, 42). In mosquito cells, sfRNA has 303 been reported as an antagonist of the RNAi response in vitro (54, 55) and contributes to 304 enhance the *in vivo* infection of mosquitoes and further dissemination from the midgut into 305 the haemocoel (32) and subsequent salivary gland infection (44). Villordo et al. (2015) 306 previously demonstrated that when passaging DENV-2 20 times in C6/36 mosquito cells, SL-307 II is highly mutated while the upstream SL-I mutates mostly upon passaging in mammalian 308 cells (40). The mutations in SL-II were shown to increase DENV-2 replication in mosquito 309 cells (40). The mutation that we found at the position 10,418 in SL-II is in line with these 310 findings, supporting the mutation pressure on SL-II in vivo, although we did not observe 311 significant mutations during passaging in C6/36 cells.

312 For DENV-2, it has been shown that during replication in human cells, mainly sfRNA1 313 is produced, while mosquito-adapted DENV-2 accumulates more abundant sfRNA3 and 314 sfRNA4 (33). We show that DENV-1 1806 produced abundant sfRNA1 while quantities of 315 sfRNA2,3 and possibly sfRNA4 were below the detection limit (Fig. 7), suggesting possible 316 differences in the production of sfRNA species between DENV-1 and DENV-2. Despite the 317 presence of the 10,418 mutation, the Ae. albopictus adapted DENV-1 1806 did not show a 318 significantly altered production of sfRNA species. Although we cannot exclude an effect on 319 cellular binding partners that might require an intact 3'UTR for their interaction with the viral 320 genome (38), it is unlikely that sfRNAs were a primary driver of DENV-1 adaptation to Ae. 321 albopictus.

We used reverse genetics to evaluate the effect of the 10,418 mutation on DENV-1 transmission by *Ae. albopictus* mosquitoes *in vivo*, but our results did not provide experimental support for a phenotypic effect of the 10,418 mutation alone. For two different

325 genetic constructs harboring the 10,418 mutation (together with different adventitious 326 mutations), there was no detectable difference in transmission efficiency. Introducing the 327 10,418 did not recapitulate the adapted phenotype of the P10 viruses and points to a more 328 complex adaptive landscape than a single-mutation effect. Because our genetic constructs 329 focused on the 10,418 mutation did not include other mutations present in the P10 viruses, it 330 implies that the enhanced transmission phenotype reflected the combined effect of several 331 mutations. Such epistatic relationships have been documented to shape the adaptive 332 landscapes of CHIKV (56, 57) and more recently DENV (58). Interestingly, the 10,418 333 mutation was the only shared mutation among replicates of adapted viruses (Fig. 3), 334 indicating that DENV-1 adaptation to Ae. albopictus can result from distinct evolutionary 335 trajectories involving different sets of mutations.

Our experimental approach has succeeded in enhancing the transmission of DENV-1 by multiple passages in the *Ae. albopictus* vector. This protocol can be extended to other arboviruses and vector species, and contribute to predict future epidemic variants. Together, this may ultimately lead to new insights into the mechanisms of arbovirus transmission by mosquitoes.

341

342 Materials and Methods

343 Ethics statement

The Institut Pasteur animal facility has received accreditation from the French Ministry of Agriculture to perform experiments on live animals in compliance with the French and European regulations on care and protection of laboratory animals (EC Directive 2010/63, French Law 2013-118, February 6th, 2013). This study was approved by the Ethics Committee #89 and registered under the reference APAFIS#6573-201606l412077987 v2. Mice were only used for mosquito rearing as a blood source, according to approved protocol.

350

351 Cell cultures

352 Ae. albopictus C6/36 cells were maintained at 28°C in Leibovitz L-15 medium supplemented 353 with non-essential amino-acids (NEAA) (1X), 10% fetal bovine serum (FBS), 100 units/mL 354 penicillin and 100 µg/mL streptomycin. These cells are defective in typical siRNAs, the 355 hallmark of exogenous RNAi mediated antiviral immunity (59). Ae. albopictus U4.4 cells 356 were maintained in L-15 medium supplemented with non-essential amino-acids (1X), 10% 357 FBS, 100 units/mL penicillin and 100 µg/mL streptomycin at 28°C. HFF (Human Foreskin Fibroblast) cells were maintained at 37°C, 5% CO2 in Dulbecco's Modified Eagle medium 358 359 (DMEM) supplemented with pyruvate, 10% FBS, 100 units/mL penicillin and 100 µg/mL 360 streptomycin. The human embryonic kidney HEK-293 cells (ATCC number CCL-1573) were 361 grown at 37°C with 5% CO2 in tissue-culture flasks with vented caps, in a minimal essential 362 medium (MEM, Life Technologies) supplemented with 7% FBS, 1% PS and 1X NEAA. 363 364 Viruses

We used two DENV-1 strains isolated from DF cases: DENV-1 1806 from an autochthonous 365 366 case from Nice, France in 2010 (provided by the National Reference Center of Arboviruses, 367 France) and DENV-1 30A from a patient in Kamphaeng Phet, Thailand in 2010 (provided by the Afrims, Thailand and under accession number HG316482 in GenBank). The 2nd passage 368 of DENV-1 1806 on African green monkey kidney Vero cells (60) and the 2nd passage of 369 370 DENV-1 30A on C6/36 Ae. albopictus cells (61) were used for mosquito infections. Serial 371 dilutions were used to determine the titer of viral stocks that was expressed in focus-forming 372 units (FFU)/mL.

373

374 Mosquito strains

375 Six populations of *Ae. albopictus* have been established from eggs: Genoa (Italy), Alessandria
376 (Italy), Cornella (Spain), Martorell (Spain), Nice Jean Archet (France), and Saint-Raphael

377 (France) (Table 1). They were tested to appraise vector competence to DENV-1 isolates. 378 Together with Ae. albopictus Nice Jean Archet (France), Ae. aegypti Pazar (Turkey) was 379 utilized to compare vector competence using viruses isolated after 10 passages on Ae. 380 albopictus. Eggs were collected from ovitraps and sent to the Institut Pasteur in Paris, where 381 they were reared in standardized conditions. After hatching, larvae were distributed in pans 382 containing a yeast tablet renewed as needed in 1 L of tap water. Adults were placed in cages 383 maintained at 28±1°C, at relative humidity of 80% and a light:dark cycle of 16h:8h, with free 384 access to 10% sucrose solution. Oral infection experiments were performed using mosquitoes 385 from the F2-F11 generations. Owing to the limited number of mosquitoes, only one biological 386 replicate was performed for each pairing population-virus.

387

388 Mosquito infections

One-week-old females were starved 24 hrs prior an infectious blood-meal in a BSL-3 laboratory. Five batches of 60 mosquito females were then allowed to feed for 15 min through a piece of pork intestine covering the base of a Hemotek feeder containing the infectious blood-meal maintained at 37°C. Only engorged females were kept and incubated under controlled conditions (28±1°C, relative humidity of 80%, light:dark cycle of 16h:8h).

394 For vector competence assays

Fourteen and 21 days after an infectious blood-meal provided at a titer of 10⁷ FFU/mL, vector competence was assessed based on two phenotypes: (i) viral infection of mosquito and (ii) viral dissemination from the midgut into mosquito general cavity. Infection rate (IR) was determined as the proportion of mosquitoes with infected midgut and dissemination efficiency (DE) was defined as the percentage of mosquitoes with virus detected in heads suggesting a successful viral dissemination from the midgut. IR and DE were calculated by titrating body and head homogenates.

402 For serial passages

403 As the first autochthonous DENV cases were reported in Nice in 2010 (9), Ae. albopictus 404 isolated in Nice was used to achieve the experimental selection of DENV-1 isolates (Fig. 2). 405 Mosquitoes were orally infected with DENV-1 supernatant provided in a blood-meal at a final titer of 10^{6.5} FFU/mL using the hemotek system. Engorged mosquitoes were incubated at 406 407 28°C for 19-21 days and then processed for saliva collection. 15-25 saliva were pooled and 408 the volume of the pool was adjusted to 600 μ L with DMEM prior to filtration through a 409 Millipore H membrane (0.22 μ m). An aliquot of 300 μ L of each sample was used to inoculate a sub-confluent flask (25 cm²) of C6/36 Ae. albopictus cells. After 1 hr, the inoculum was 410 411 discarded and cells were rinsed once with medium. Five mL of DMEM medium 412 complemented with 2% FBS was added and cells were incubated for 8 days at 28°C. Cell 413 culture supernatants were then collected and provided to mosquitoes to run the next passage. 414 Passages P1 to P3 were performed with mosquitoes of the F3 generation and passages P4 to 415 P10 with mosquitoes of the F4 generation. C6/36 supernatants collected at each passage were 416 used undiluted for the next mosquito blood-meal. Ten passages were performed. Control 417 isolates corresponded to serially passaged viruses on C6/36 cells to identify mutations 418 resulting from genetic drift or adaptation to insect cell line; 500 μ L of the previous passage 419 were used to inoculate the next flask of C6/36 cells. Two biological replicates R1 and R2 420 were performed to test the variability between samples submitted to the same protocol of 421 selection. Vector competence using the parental and P10 isolates was assessed by calculating: 422 (i) infection rate (IR, proportion of mosquitoes with infected midgut), (ii) dissemination 423 efficiency (DE, proportion of mosquitoes able to disseminate the virus from the midgut 424 among tested mosquitoes), and (iii) transmission efficiency (TE, proportion of mosquitoes 425 with the virus detected in saliva among tested mosquitoes).

426

427 Virus deep sequencing

428 Total RNA was extracted from cell culture supernatant using QIAamp Viral RNA Mini Kit 429 (Qiagen, Germany) and DNAse treated (Turbo DNAse, Life Technologies, USA). Following 430 purification with magnetic beads (Agencourt RNAClean XP, Beckman Coulter, California, 431 USA), RNA was reverse transcribed using Transcriptor High Fidelity cDNA Synthesis Kit 432 and a specific 3'-UTR DENV-1 primer (Roche Applied Science, Mannheim, Germany), 433 d1a5B 5'-AGAACCTGTTGATTCAACRGC-3' (62). Second strand was then synthetized in 434 a unique reaction with E. coli DNA ligase (New England Biolabs, Massachusetts, USA), E. 435 coli DNA polymerase I (New England Biolabs), E. coli RNAse H (New England Biolabs) in 436 second strand synthesis buffer (New England Biolabs). After purification with magnetic beads 437 (Agencourt AMPure XP, Beckman Coulter), dsDNA was quantified with fluorometric 438 method (Quant-iT PicoGreen dsDNA, Invitrogen, Massachusetts, USA).

439 Sequencing libraries were prepared using Nextera XT DNA Library Preparation Kit 440 (Illumina, San Diego, USA), multiplexed and sequenced in single end in two independent 441 runs on an Illumina NextSeq 500 platform using a mid-output 150-cycle v2 kit (Illumina). 442 Reads were trimmed (Trimmomatic v0.33) (63) after demultiplexing (bcl2fastq v.2.15.0, 443 Illumina) to remove adaptor sequences, and reads shorter than 32 nucleotides were discarded. 444 Full-length genome of the DENV-1 1806 was assembled *de novo* using Ray v2.0.0 (64) with 445 the original stock sample. The newly assembled DENV genome contig was extended in 3' 446 and 5' using closest BLAST hit full DENV-1 genome (accession number EU482591). This 447 chimeric construct was used to map reads used for assembly using Bowtie 2 v2.1.0 (65). 448 Alignment file was converted, sorted and indexed using Samtools v0.1.19 (66). Coverage and 449 sequencing depth were assessed using bedtools v2.17.0 (67). Single nucleotide variants and 450 their frequency were called using LoFreq* v2.1.1 (68) and used to correct the chimeric 451 construct. Only nucleotides with >10X coverage were conserved for generating the consensus 452 sequence. A final full-length genome sequence for DENV-1 1806 strain was deposited to 453 GenBank (accession number MG518567).

After quality control, reads from all samples were mapped to the newly assembled DENV-1 1806 strain genome sequence or previously sequenced reference genome KDH0030A (accession number HG316482) using Bowtie v2.1.0 (65). The alignment file was converted, sorted and indexed using Samtools v0.1.19 (66), and the coverage and sequencing depth were assessed for each sample using bedtools v2.17.0 (67). Single nucleotide variants (SNVs) and their frequency were then called using LoFreq* v2.1.1 (68), and their effect at the amino-acid level was assessed by SNPgenie v1.2 (69).

461

462 **RNA structure modeling in silico**

The Mfold Web server was used with standard settings and flat exterior loop type (70) to fold the secondary RNA structures, which were then visualized using the VARNA RNA editing package (71). Pseudoknot RNA interactions were drawn as previously described for DENV (45, 72). Mutation frequencies of individual nucleotides were determined by averaging the nucleotide allele frequency from the deep sequencing results of the duplicates per treatment.

468

469 Virus growth curves

470 To measure viral replicative fitness, growth curves were conducted in Ae. albopictus C6/36 471 and U4.4 mosquito cells, and Human Foreskin Fibroblasts (HFF) cells. Confluent cell 472 monolayers were prepared and inoculated with viruses simultaneously in triplicates at a MOI 473 of 0.1 PFU/cell. Cells were incubated for 1 hr in appropriate conditions and viral inoculum 474 was removed to eliminate free virus. Five mL of medium supplemented with 2% FBS were 475 then added and mosquito cells were incubated at 28°C (mosquito cells) or 37°C (human 476 cells). At various times (4, 6, 8, 10, 24, 48 and 72 hrs) post-inoculation (pi), supernatants were 477 collected and titrated by focus fluorescent assay on Ae. albopictus C6/36 cells. After 478 incubation at 28°C for 5 days, plates were stained using hyper immune ascetic fluid specific 479 to DENV as primary antibody (Millipore, Molsheim, France). A Fluorescein-conjugated goat

anti-mouse was used as the second antibody (Thermofisher). Three viral strains were used:
the parental strain and two 10th passages, P10_R1 and P10_R2. Viral titer was expressed in
FFU/mL. Three biological replicates were performed for each cell-virus pairing.

483

484 RNA isolation and Northern blotting

485 Total RNA was isolated from cell monolayers using TRIzol reagent (Invitrogen, 486 Massachusetts, France) following the manufacturer's protocol. Mosquito DENV-1 infected 487 bodies were homogenized individually in 500 µL of Leibovitz L15 medium (Invitrogen) 488 supplemented with 2% fetal bovine serum for 1 min at maximum speed. Homogenates were 489 then filtered with a filter unit (0.22 µm) (Ultrafree® MC-GV, Merck, New Jersey, USA). 490 Two samples of each filtrate were inoculated onto monolayers of Ae. albopictus C6/36 cell 491 culture in 6-well plates. After incubation at 28°C for 6 days, samples were homogenized with 492 1 mL TRIzol reagent. RNA isolations were performed using the standard TRIzol protocol. 493 Samples were eluted in 30 µL RNase-free Milli-Q water and stored at -80°C until further 494 processing. A DENV-1 3'UTR specific probe was generated by PCR reaction with GoTaq 495 Polymerase (Promega, Wisconsin, USA) containing DIG DNA-labelling mix (Roche) and 496 primers DENV-1 3'UTR FW (AGTCAGGCCAGATTAAGCCATAGTACGG) and DENV-1 497 3'UTR RV (ATTCCATTTTCTGGCGTTCTGTGCCTGG) using cDNA from cells infected 498 with DENV-1 1806 as a template. Five micrograms of total RNA was subjected to sfRNA-499 optimized northern blot as has been described previously (32). Briefly, total RNA was 500 denatured and size separated on 6% polyacrylamide-7 M urea-0.5× Tris-borate-EDTA (TBE) 501 gel for 1.45 hrs at 150 V. The RNA was semi-dry-blotted on a Hybond-N membrane, UV 502 cross-linked and pre-hybridized for 1 hr at 50°C in modified Church buffer containing 10% 503 formamide. DENV-1 3'UTR specific Dig-labelled probe was denatured and blots were 504 hybridized overnight at 50°C in modified church/10% formamide buffer containing 2 μ L of 505 DIG-labelled probe. Blots were developed with AP-labeled anti-DIG antibodies and NBT-

BCIP solution before observing the signal using a Bio-Rad Gel Doc scanner. Quantification of band signal intensities was performed in ImageJ by transforming the image to 8-bit format, inverting the image, and analyzing the band intensity using the measure function. The Ratio sfRNA/gRNA was calculated by dividing the intensity of the sfRNA by the intensity of the gRNA band for each sample, and then normalized to the average ratio of the parental samples.

511

512 **ISA reverse genetics**

The T>C mutation at position 10,418 identified at passage 10 was inserted into a DENV-1 1806 backbone using the ISA (Infectious Subgenomic Amplicons) reverse genetics method as previously described (73).

516 Preparation of subgenomic DNA fragments

517 The viral genome was amplified by RT-PCR from the DENV-1 1806 viral RNA as three 518 overlapping DNA fragments. Two additional fragments were *de novo* synthesized (Genscript) 519 and amplified by PCR (primers are listed in Table S6). The first primer consisted of the 520 human cytomegalovirus promoter (pCMV) and the second primer of the last 367 nucleotides 521 of the 3'UTR of the DENV-1 1806 with or without the 10,418 T>C mutation and the hepatitis 522 delta ribozyme followed by the simian virus 40 polyadenylation signal (HDR/SV40pA) 523 (sequences are listed in Supplemental Text). RT mixes were prepared using the superscript IV reverse transcriptase kit (Life Technologies, CA, USA) and PCR mixes using the Q5® High-524 Fidelity PCR Kit (New England Biolabs, MA, USA) following the manufacturer's 525 526 instructions. RT were performed in the following conditions: 25° C for 10^{\Box} min followed by 527 37□°C for 50 min and 70°C 15□min. PCR amplifications were performed in the following 528 conditions: $98 \square ^{\circ}C$ for $30 \square$ sec followed by 35 cycles of $98 \square ^{\circ}C$ for 10 s, $62 \square ^{\circ}C$ for $30 \square$ s, 529 $72 \square ^{\circ}C$ for $2 \square \min 30$ s, with a 2 min final elongation at $72 \square ^{\circ}C$. PCR product sizes and 530 quality were controlled by running gel electrophoresis and DNA fragments were purified 531 using a QIAquick PCR Purification Kit (Qiagen, Hilden, Germany).

532 *Cell transfection*

533 HEK-293 cells were seeded into six-well cell culture plates one day prior to transfection. 534 Cells were transfected with 2 µg of an equimolar mix of the five DNA fragments using 535 lipofectamine 3000 (Life Technologies) following the manufacturer's instructions. Each 536 transfection was performed in five replicates. After incubating for 24 hrs, the cell supernatant 537 medium was removed and replaced by fresh cell culture medium. Seven days post-538 transfection, cell supernatant medium was passaged two times using six-well cell culture 539 plates of confluent C6/36 cells. Cells were subsequently inoculated with 100 μ L of diluted 540 (1/3) cell supernatant media, incubated 1 hr, washed with PBS 1X, and incubated 7 days with 541 3 mL of medium. Remaining cell supernatant medium was stored at $-80 \square$ °C. The second 542 passage was used to produce virus stock solutions of DENV-1 1806 WT and mutant viruses. 543 Transmission efficiency was assessed 21 days after an infectious blood meal containing the 544 Parental, the Parental construct, the P10 strain, the P10 constructs (1 and 2) provided separately at a titer of 10^7 FFU/mL. 545

546

547 **Statistical analyses**

Statistical analyses were conducted using the STATA software (StataCorp LP, Texas, and
USA). P-values>0.05 were considered non-significant. If necessary, the significance level of
each test was adjusted based on the number of tests run, according to the sequential method of
Bonferroni (74).

552

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568

569 Author contributions

- 570 LL and ABF designed the research. RB, SL, HJ, GPG, LM, GP, and FA performed
- 571 experiments. LM, GP, PSY, GG, and MV provided reagents and analytical tools. AS and
- 572 XDL provided expertise and feedback. RB, SL, HJ, GPG, GPP, and ABF analyzed the data.
- 573 GPG, GPP, LL, and ABF wrote the manuscript. LL and ABF secure funding.

574

575 **Declaration of interests**

- 576 The authors declare no competing interests.
- 577

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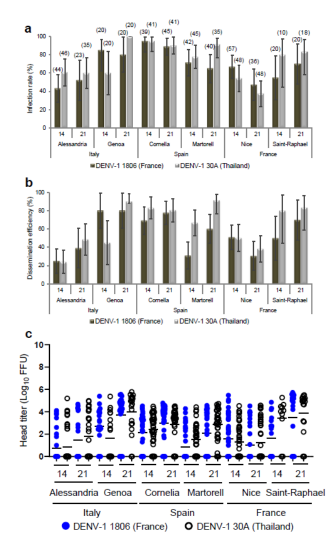
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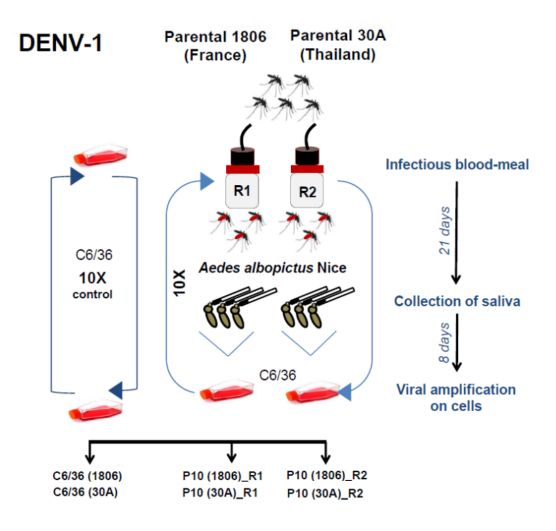
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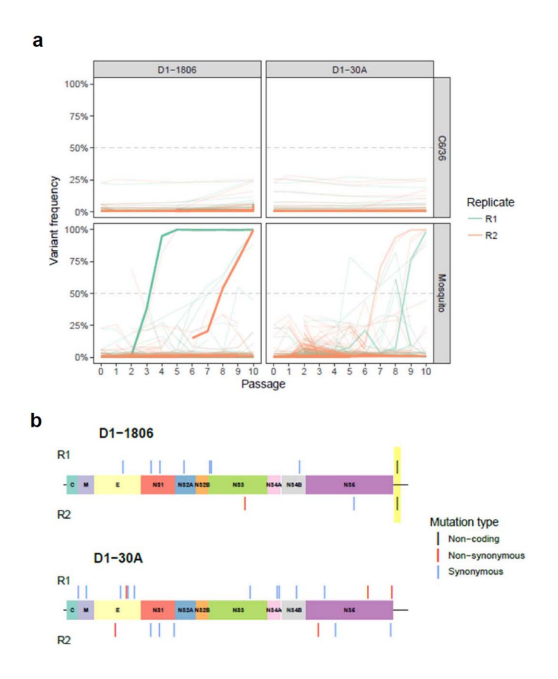
769 Figure 1. Susceptibilities of six European Ae. albopictus populations to DENV-1 (1806 and 30A): **a** infection rate, **b** dissemination efficiency and **c** viral titers in heads. Adult female 770 mosquitoes were challenged with DENV-1 from France (1806) and Thailand (30A) at a titer 771 of 10⁷ FFU/mL. At 14 and 21 dpi, mosquitoes were sacrificed and decapitated. Bodies and 772 heads were homogenized and titrated on C6/36 cells. Infection rates were determined using 773 774 positive/negative scoring (i.e. without estimating the number of viral particles), while viral 775 titers at sites of dissemination were quantified via focus-forming assay. The error bars 776 correspond to the confidence intervals (95%) (\mathbf{a}, \mathbf{b}) , and the bar to the mean (\mathbf{c}) . In brackets is 777 indicated the sample size.



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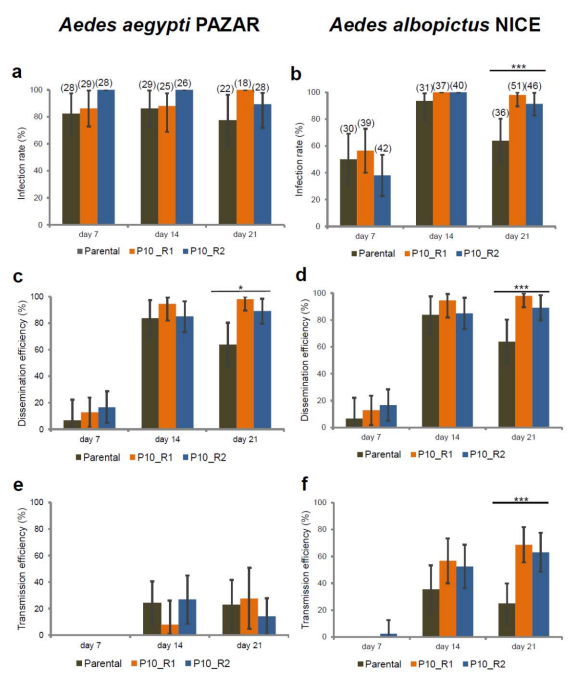
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Figure 2. Experimental design for DENV-1 adaptation to *Ae. albopictus*. The parental strains 1806 (France) and 30A (Thailand) were passaged 10 times on a single *Ae. albopictus* population from Nice, France. Each passage includes: mosquito infectious blood-meal with DENV-1, collection of mosquito saliva at day 21 post-infection, viral amplification of saliva on *Ae. albopictus* cell cultures for 8 days, and initiation of the next passage using the viral suspension obtained. Control isolates were serially passaged 10 times on C6/36 cells. Two replicates R1 and R2 were performed.



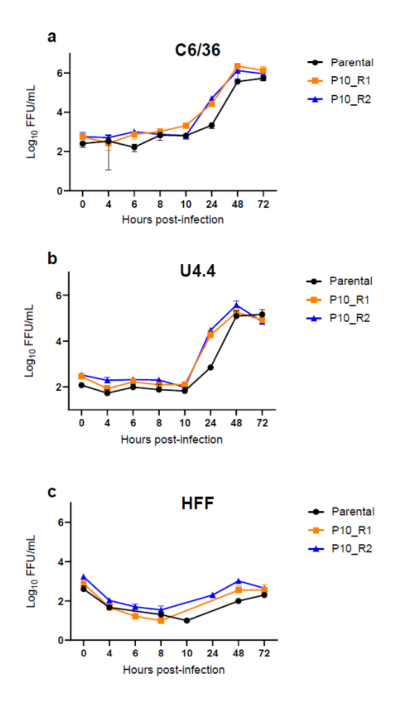
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Figure 3. Frequency variation and genomic position of consensus level-reaching variants during passages. a The top panels correspond to DENV-1 passaged on C6/36 cells as controls and the bottom panels to DENV-1 passaged on *Ae. albopictus* mosquitoes. Bold lines represent the variant 10,418 in the 3'UTR. b Variants are represented with a colored segment according to the mutation type (non-coding: black; non-synonymous: red; synonymous: blue). The position of the only shared variant between two replicates is highlighted in yellow.



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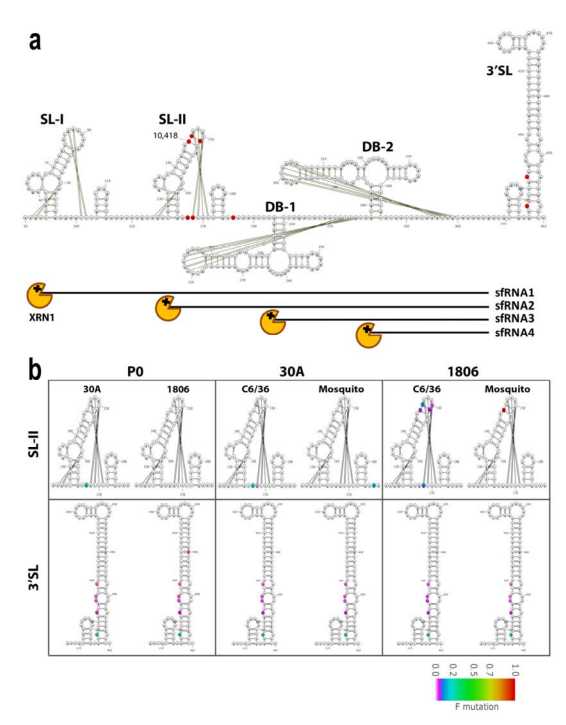
Figure 4. Infection, Dissemination and Transmission of DENV-1 (Parental, P10_R1, and 799 P10_R2) by Ae. aegypti Pazar and Ae. albopictus Nice. Mosquitoes were exposed to blood 800 801 meals at a titer of 10^{7} FFU/mL. Females were examined at 7, 14 and 21 dpi. Mosquito body (thorax and abdomen) and head were processed individually to determine (a, b) the infection 802 803 rate (IR, proportion of mosquitoes with infected body among the engorged mosquitoes) and 804 (c, d) the dissemination efficiency (DE, proportion of mosquitoes with infected head among 805 tested mosquitoes). (e, f) Saliva was collected from individual females to determine the transmission efficiency (TE, proportion of mosquitoes with infectious saliva among tested 806 807 mosquitoes). The Parental strain corresponds to DENV-1 1806, and P10_R1 and P10_R2 refer, respectively, to replicate 1 and replicate 2 of the 10th in vivo passages of DENV-1 1806 808 on Ae. albopictus. Asterisks refer to a significant difference (*** $p < 10^{-3}$). In brackets, the 809 810 number of mosquitoes tested. The error bars correspond to the confidence intervals (95%). 811



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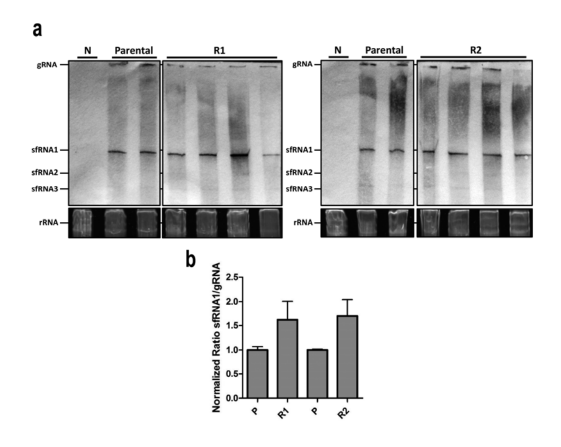
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Figure 5. Growth curves of the two passages 10 of DENV-1 1806 strain in two cell lines, (**a**) *Ae. albopictus* C6/36 cells, (**b**) *Ae. albopictus* U4.4 cells, and (**c**) human foreskin fibroblasts HFF cells. Cells were infected with the parental strain and the two replicates of the 10th passages of DENV-1 1806 (P10_R1 and P10_R2) at a MOI of 0.1. Supernatants were collected at 4, 6, 8, 10, 24, 48 and 72 hrs post-inoculation. The number of infectious viral particles was determined by focus fluorescent assay on *Ae. albopictus* C6/36 cells. Three replicates were performed for each cell-virus pairing. Error bars show standard deviations.



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- 823

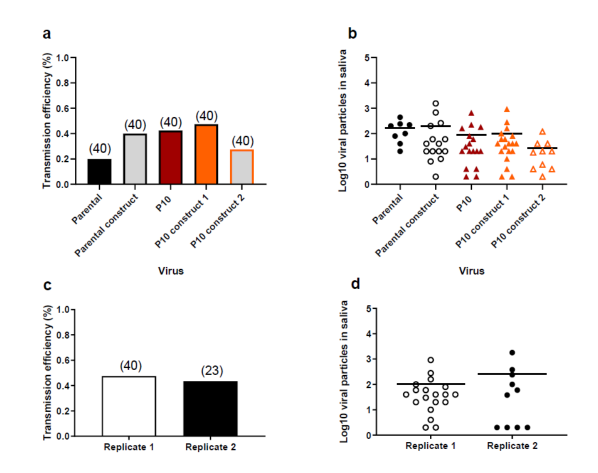
824 Figure 6. Analysis of mutation frequencies in the DENV-1 3'UTR. a Schematic overview of 825 the DENV-1 3'UTR secondary RNA structure, indicating from 5' to 3' the stem loop (SL)-I, 826 SL-II, dumbbell (DB)-1, DB-2 and 3'SL RNA structures. Single nucleotide variants with a 827 frequency ≥0.05 after 10 passages of DENV-1 1806 or 30A in either C6/36 or mosquitoes are 828 highlighted in red. Pseudoknots and other tertiary RNA interactions are indicated by the black 829 lines. **b** Analysis of the mutation frequencies in SL-II and the 3'SL of the parental DENV-1 830 30A and 1806 sequences, and passages P10 (1806) and P10 (30A). The mutation frequency is 831 indicated by color on a scale from 0 (white) to 1 (red). 832



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835 Figure 7. Northern blot detection of sfRNA production after C6/36 cells infection with 836 DENV-1 1806. a Visualization of sfRNA. 5µg of total RNA from the parental and the two 837 replicates P10 R1 and P10 R2 of DENV-1 1806 or non-infected cells (N) was size separated on a 6% Polyacrylamide/Urea gel. One gel was run for the R1 samples (left) and one for the 838 839 R2 samples (right), including the negative and parental samples on each gel. Then, RNAs 840 were blotted onto Hybond-N paper and subjected to northern-blotting with a DENV-1 3'UTR 841 specific probe. The bands shown correspond to the DENV genomic RNA (gRNA) and subgenomic flavivirus RNA (sfRNA). As loading control, the ribosomal RNA (rRNA) from 842 843 the EtBr stained gel are shown. **b** Quantification of the ratio of sfRNA to gRNA production. 844 Band intensities were determined using the 'Measure' function in ImageJ. The intensity of the 845 background (lane N) was subtracted from the readings before the ratio sfRNA/gRNA was 846 calculated by dividing the intensity of the sfRNA by the intensity of the gRNA band for each 847 sample, and then normalized to the average ratio of the parental samples. The statistics were 848 performed using a two-tailed unpaired t-test.



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852 Figure 8. Reverse genetic constructs with the 10,418 mutation do not show higher transmission in Ae. albopictus. a Transmission of reverse genetic constructs (Parental 853 construct, P10 construct 1, P10 construct 2) by Ae. albopictus Nice with reference to Parental 854 and P10 strains. Twenty-one days after an infectious blood meal at a titer of 10^7 FFU/mL, 855 mosquitoes were processed for saliva collection to determine the transmission efficiency (TE, 856 857 proportion of mosquitoes with infectious saliva among tested mosquitoes). b Viral loads in 858 saliva were estimated by focus fluorescent assay on Ae. albopictus C6/36 cells. c, d A second 859 replicate using the P10 construct 1 was performed. Bars indicate the mean. 860

Table 1. Details on mosquito populations used for experimental infections with DENV-1.

Species	Mosquito population	Country	Date of collection	Generation used for mosquito infections		
	Nice	France	August 2011	F3, F11		
	Saint-Raphaël	France	November 2012	F3		
Aedes	Alessandria	Italy November 2012		F1		
albopictus	Genoa	Italy	October 2012	F2		
	Cornella	Spain	July 2012	F2		
	Martorell	Spain	July 2012	F2		
Aedes aegypti	Pazar	Turkey	2016	F4		

Table S1. Comparisons of infection rates and dissemination efficiencies between mosquitoes

- see infected with two DENV-1 strains (1806 and 30A) and examined at different days post-
- 867 infection (14 and 21).

		Fisher's exact test			
Mosquito population	Day post-infection	Infection rate	Dissemination efficiency		
Alessandria	14	0.093	0.905		
Alessandria	21	0.556	0.479		
Genoa	14	0.077	0.022		
Genoa	21	0.035	0.376		
Cornella	14	0.979	0.168		
Comena	21	0.808	0.715		
Martorell	14	0.496	0.001		
Wattoren	21	0.006	0.002		
Nice	14	0.191	0.929		
Nice	21	0.371	0.463		
Saint-Raphael	14	0.18	0.114		
Samercaphaer	21	0.334	0.334		

- 871 Table S2. Comparisons of infection rates and dissemination efficiencies between the 6
- mosquito populations infected with a DENV-1 strain (1806 or 30A) and examined at a given
- 873 day post-infection (14 or 21).

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	Day post-infection	Fisher's exact test			
DENV-1		Infection rate	Dissemination		
		Intection rate	efficiency		
1806	14	0.0001	0.0001		
	21	0.001	0.0001		
30A	14	0.0001	0.0001		
	21	0.0001	0.0001		

875

- 877 Table S3. Comparisons of viral loads in mosquito heads between mosquitoes infected with
- two DENV-1 strains (1806 and 30A) and examined at different days post-infection (14 and
- 879 21).
- 880

Mosquito population	Day post-infection	Wilcoxon Rank-Sum test
Alessandria	14	0.292
Alessalulla	21	0.746
Genoa	14	0.533
Genoa	21	0.836
Cornella	14	0.439
Comena	21	0.018
Martorell	14	0.073
Wattoren	21	0.304
Nice	14	0.183
TVICC	21	0.857
Saint-Raphael	14	0.032
Samt-Kaphaci	21	0.169

bioRxiv preprint doi: https://doi.org/10.1101/2020.07.16.206524; this version posted July 17, 2020. The copyright holder for this preprint (which was not certified by peer review) is the author/funder, who has granted bioRxiv a license to display the preprint in perpetuity. It is made available under aCC-BY-NC-ND 4.0 International license. **Table S4.** Position and estimated frequency of SNVs reaching consensus level detected in the

mosquito-passaged samples.

Virus	Passage	Replicate	SNV position on DENV genome	Reference nucleotide	SNV	Sequencing depthat position	S NV frequency	Position incodon	Mutation type
1806	2	2	5667	Т	с	14231	0,686319	2	Non-synonymo
1806	4	1	4606	А	G	7484	0,953234	3	Sy no ny mo us
1806	4	1	73 60	Т	С	5336	0,962331	3	Synonymous
1806	4	1	10418	Ť	c	3703	0,94734	NA	non-coding
1806	4	2	9067	T	С	3998	0,815408	3	Sy no ny mo us
1806	5	1	4606	A	G	141288	0,999023	3	Sy no ny mo us
1806	5	1	73 60	Т	с	29 2005	0,999017	3	Sy no ny mo us
1806	5	1	10418	Т	С	199563	0,998943	NA	non-coding
1806	5	2	9067	T	с	94413	0,998856	3	Synonymous
1806	6	1	4606	A	G	37980	0,996288	3	Sy no ny mo us
1806	6	1	7360	Т	С	20048	0,996758	3	Sy no ny mo us
1806	6	1	10418	Т	С	18264	0,996605	NA	non-coding
1806	6	2	9067	Т	с	21252	0,996518	3	Sy no ny mo us
1806	7	1	3757	С	T	16059	0,55869	3	Sy no ny mo us
1806	7	1	4606	A		24880		3	
					G		0,996383		Sy no ny mo us
1806	7	1	73 60	T	С	15019	0,997004	3	Sy no ny mo us
1806	7	1	10418	T	с	9 195	0,996194	NA	non-coding
1806	7	2	9067	т	С	5095	0,995682	3	Sy no ny mo us
1806	8	1	27 19	А	G	7900	0,553544	3	Sy no ny mo us
	8	1		с	T	8388		3	
1806			3001				0,510968		Sy no ny mo us
1806	8	1	3757	С	T	9 649	0,643694	3	Sy no ny mo us
1806	8	1	4552	G	А	10485	0,623176	3	Sy no ny mo us
1806	8	1	4606	А	G	11782	0,996605	3	Sy no ny mo us
1806	8	1	73 60	T	c	10285	0,996889	3	Synonymous
1806	8	1	10418	T	c	6829	0,997218	NA	non-coding
1806	8	2	9067	T	С	4272	0,994616	3	Sy no ny mo us
1806	8	2	10418	T	С	3477	0,540696	NA	non-coding
1806	9	1	1840	С	Т	16264	0,545315	3	Sy no ny mo us
1806	9	1	27 19	А	G	14188	0,790668	3	Sy no ny mo us
1806	9	1	3001	c	 Т	12951	0,792448	3	Syno ny mo us
1806	9	1	3757	с	T	15968	0,830348	3	Sy no ny mo us
1806	9	1	4552	G	А	15493	0,828632	3	Sy no ny mo us
1806	9	1	4606	A	G	17632	0,996257	3	Sy no ny mo us
1806	9	1	73 60	Т	с	15973	0,997308	3	Sy no ny mo us
1806	9	1	10418	T	С	10027	0,995412	NA	non-coding
1806	9	2	9067	T	С	6528	0,996477	3	Sy no ny mo us
1806	9	2	10418	Т	С	5922	0,761905	NA	non-coding
1806	10	1	27 19	A	G	84568	0,877956	3	Sy no ny mo us
1806	10	1	3001	с	T	7 2183	0,972334	3	Sy no ny mo us
1806	10	1	3757	С	T	9 27 00	0,975955	3	Synonymous
1806	10	1	4552	G	A	101912	0,974851	3	Sy no ny mo us
1806	10	1	4606	A	G	101144	0,9991	3	Sy no ny mo us
1806	10	1	73 60	Т	с	213796	0,998948	3	Sy no ny mo us
1806	10	1	10418	Т	с	141416	0,998883	NA	non-coding
1806	10	2	9067	Т	С	29741	0,998352	3	Synonymous
1806	10	2	10418	Ť	c	37209	0,995996	NA	
									non-coding
30A	5	1	1768	T	A	312358	0,787503	3	Sy no ny mo us
30A	6	1	1768	Т	A	1264	0,665348	3	Sy no ny mo us
30A	7	1	1768	Т	A	8 2 9 0	0,563209	3	Sy no ny mo us
30A	7	2	1611	С	Т	8758	0,704042	2	Non-synonymo
30A	7	2	2977	T	с	9778	0,710268	3	Synonymous
30A	7	2	3442	G	А	5856	0,691257	3	Sy no ny mo us
30A	7	2	10208	С	T	5068	0,608327	1	Sy no ny mo us
30A	8	1	448	Т	с	1075	0,6	3	Sy no ny mo us
30A	8	1	1768	Т	А	2389	0.821683	3	Sy no ny mo us
30A	8	1	1959	A	G	2247	0,604806	2	Non-synonymo
							0,519898		
30A	8	1	5822	C T	T	1181		1	Sy no ny mo us
30A	8	1	8149	T	С	1248	0,615385	3	Sy no ny mo us
30A	8	1	10258	A	Т	1017	0,612586	3	Non-synonymo
30A	8	2	1611	С	Т	5281	0,937512	2	Non-synonymo
30A	8	2	2977	Т	c	5 19 2	0,943374	3	Sy no ny mo us
30A 30A	8	2	3442	G	A	3062	0,943374	3	Synonymous
30A	8	2	10208	с	T	1783	0,902973	1	Sy no ny mo us
30A	9	1	694	С	T	1501	0,774151	3	Sy no ny mo us
30A	9	1	2002	с	Т	2058	0,757532	3	Sy no ny mo us
30A	9	1	6658	G	А	1778	0,72216	3	Sy no ny mo us
30A	9	1	67 28	c	T	2246	0,769813	1	Synonymous
	9	1		С	T				
30A			7267			978	0,747444	3	Sy no ny mo us
30A	9	1	9504	A	G	1527	0,762279	2	Non-synonymo
30 A	9	2	1611	С	T	11211	0,997681	2	Non-synonymo
30 A	9	2	27 16	A	G	5452	0,762656	3	Sy no ny mo us
30A	9	2	2977	T	С	11160	0,996416	3	Sy no ny mo us
30A	9	2	3442	G	A	7394	0,994725	3	
									Synonymous
30A	9	2	7952	T	с	11955	0,931409	1	Non-synonymo
30A	9	2	8485	T	С	13405	0,898396	3	Sy no ny mo us
30A	9	2	10208	С	T	5542	0,997835	1	Sy no ny mo us
30A	10	1	694	С	Т	199 24	0,984039	3	Sy no ny mo us
30A	10	1	2002	c	T	35756	0,984758	3	Sy no ny mo us
30A	10	1	2200	T	с	20712	0,876014	3	Sy no ny mo us
30A	10	1	6658	G	А	18057	0,976574	3	Sy no ny mo us
	10	1	67 28	С	Т	23549	0,982717	1	Sy no ny mo us
30A	10	1	7267	c	T	17245	0,985503	3	Sy no ny mo us
30A 30A									
30A	10	1	9504	A	G	22552	0,982973	2	Non-synonymo
30A 30A			1611	С	T	18775	0,998189	2	Non-synonymo
30A	10	2							
30A 30A		2	2716	A	G	13145	0,909623	3	Sy no ny mo us
30A 30A 30A 30A	10 10	2	27 16						
30A 30A 30A 30A 30A 30A	10 10 10	2	27 16 29 77	Т	С	135 26	0,998743	3	Sy no ny mo us
30A 30A 30A 30A 30A 30A 30A	10 10 10 10	2 2 2	27 16 2977 3442	T G	C A	135 26 8885	0,998743 0,996511	3	Sy no ny mo us
30A 30A 30A 30A 30A 30A	10 10 10	2	27 16 29 77	Т	С	135 26	0,998743	3	Sy no ny mo us

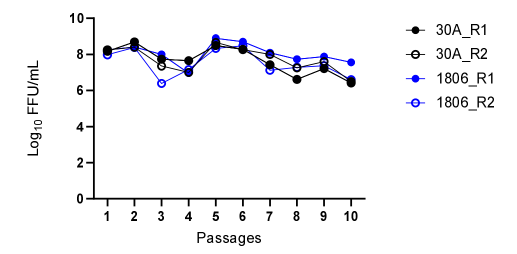
Table S5. Sequences of the three reverse genetic constructs (Parental, P10 construct 1, P10 construct 2) using the deep sequencing method described in materials and methods. Only substitutions with a mutation frequency > 5% were considered significant for further analysis. The parental strain P0 was used as the reference sequence. The two P10 constructs presented the mutation 10,418 (in grey) at a frequency close to 100% beside other mutations.

Clone	Mutation	Position	Frequency (%)	AA Position	AA change	Gene
	A->G	211	27.2	39	Ser	Capsid
Parental IC	T->C	4697	25.9	60	Tyr -> His	nsp3
Falentai IC	C->T	8650	30	359	Asp	nsp5
	T->C	9172	6.3	533	Asp	nsp5
	T->C	1513	5	193	Phe	Env
	G->T	2684	5.4	89	Val -> Phe	nsp1
	T->C	3472	5.3	351	Ser	nsp1
	A->T	3581	5.6	36	Thr -> Ser	nsp2a
P10 IC1	T->C	5173	52.5	218	Arg	nsp3
	G->A	5764	5.8	415	Gly	nsp3
	C->A	6346	5.5	609	Arg	nsp3
	A->G	7210	33.1	128	Ala	nsp4b
	T->C	10418	100			
	T->C	7321	98.8	165	Phe	nsp4b
P10 IC2	G->A	9571	98.4	666	Val	nsp5
	T->C	10418	99.9			

pCMV_F	TCAATATTGGCCATTAGCCATATTAT		
pCMV_tag_R	GTTCTTGTCGGTCCACGTAGACTAACAACTCGGTTCACTAAACGAGCTCTGC	pCMV tagged	
1806_I_tag_F	CTATATAAGCAGAGCTCGTTTAGTGAACCGAGTTGTTAGTCTACGTGGACCGA	Fragment I	
1806_I_R	CTGACCCTGCAGAGACCATTGA	tagged	
1806_II_F	1806_II_F CCACAACAGTCACAGGAAAGATA		
1806_II_R	CTCCACGTCCATGTTCTCCTC	Fragment II	
1806_III_F	CAGCAATAGACGGGGAGTACAG	Fragment III	
1806_III_WT_R	CTCAATCCGTGGCTTTCGGC	WT	
1806_III_F	CAGCAATAGACGGGGAGTACAG	Fragment III	
1806_III_Mut_R	CTCAGTCCGTGGCTTTCGGC	Mut	
<u>3UTRHDRSV40 F</u>	F GTGAGCCCCGTCCAAGGACGT		
<u>3UTRHDRSV40 R</u>	CTCAGGGTCAATGCCAGCGCT	<u>3'UTR +</u> <u>HDR/SV40pA</u> (WT and Mut)	

Table S6. Primers used to generate subgenomic DNA fragments (ISA procedure)

Figure S1. Viral titers of cell culture supernatants used to run passages in the experimental selection for DENV-1 adaptation to *Ae. albopictus*. Saliva were collected from 15-25 mosquitoes 19-21 days after infection and pooled to inoculate a monolayer of C6/36 *Ae. albopictus* cells. After 8 days at 28°C, cell culture supernatants were collected and provided to mosquitoes to run the next passage. Ten passages were performed. The supernatants were titrated by focus fluorescent assay on *Ae. albopictus* C6/36 cells. Viral titer was expressed in FFU/mL. Two biological replicates were performed for each viral strain, DENV-1 30A and DENV-1 1806.



1-1806-P0-R2-0	1-1806-P1-R1-0	-1806-P1-R1-m	1-1806-P1-R2-0	-1806-P1-R2-m	-1806-P10-R1-	-1806-P10-R1-n	I-1806-P10-R2
-1806-P10-R2-n	806-P10-R2-mo	-1806-P2-R1-m	-1806-P2-R2-m	-1806-P3-R1-m	-1806-P3-R2-m	-1806-P4-R1-m	-1806-P4-R2-I
2-1 	-1806-P5-R1-m	1-1806-P5-R2-(-1806-P5-R2-m	-1806-P6-R1-п маламала	-1806-Р6-R2-л	-1806-P7-R1-m	-1806-P7-R2-
-1806-P8-R1-m	-1806-P8-R2-m	-1806-P9-R1-m	-1806-P9-R2-m	11-30A-P0-R2-c)1-30A-P1-R1-c	1-30A-P1-R1-m)1-30A-P1-R2-
1-30A-P1-R2-m	1-30A-P10-R1-(-30A-P10-R1-m	1-30A-P10-R2-0	-30A-P10-R2-m	 1-30A-P2-R1-m		1-30A-P3-R1-r
1-30A-P3-R2-m	 -30A-P4-R1-m	 -30A-P4-R2-m	1-30A-P5-R1-c	1-30A-P5-R1-m	1-30A-P5-R2-c	1-30A-P5-R2-m	1-30A-P6-R1-r
1-30A-P6-R2-m	1-30A-P7-R1-m	1-30A-P7-R2-m	1-30A-P8-R1-m	1-30A-P8-R2-m	1-30A-P9-R1-m	1-30A-P9-R2-m	0 300060009000
	0 300060009000	ó 300060009000	0 300060009000 Position on D		0 30006000000	0 300060009000	

Figure S2. Sequencing coverage and depth by sample.

Position on DEINV genome

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Supplemental Text. Sequences of the *de novo* subgenomic DNA fragments used during the ISA procedure

<u>pCMV</u>

367 last nucleotides of 3'UTR + HDR/SV40pA (WT; T)

367 last nucleotides of 3'UTR + HDR/SV40pA (Mutant; C)