

1 Experimentally mimicking 30 years of
2 *Magallana gigas* infections with the OsHV-
3 1 virus reveals evolution through positive
4 selection

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23 Abstract

24 Ostreid herpesvirus 1 (OsHV-1) poses a significant threat to the global oyster farming industry, causing
25 substantial economic losses due to mortality outbreaks. While OsHV-1 primarily affects the Pacific
26 oyster *Magallana gigas*, it has been linked to mortality events in various host species. Despite
27 advancements in understanding OsHV-1 epidemiology, knowledge gaps persist regarding its
28 evolutionary mechanisms and adaptation to host genetic backgrounds. This study employs experimental
29 evolution and extensive genomic analysis to unravel the dynamics of OsHV-1 evolution in response to
30 oyster host genetic variation. Our results show that genetic mutations, particularly transitions and
31 transversions, played a significant role in shaping the viral population, leading to a trend toward genetic
32 homogenization. Stronger positive selection signals were observed in the oyster population with higher
33 susceptibility, suggesting adaptation of viral genotypes to specific host genetic backgrounds. These
34 findings shed light on the complex evolutionary dynamics of OsHV-1 and its interactions with oyster
35 hosts. Understanding how this virus adapts to host genetic diversity is crucial for developing strategies
36 to mitigate its impact on the oyster farming industry and provides valuable insights into the broader
37 mechanisms of viral evolution in response to host variation.

38 Introduction

39 Ostreid herpesvirus 1 (OsHV-1), a double-stranded DNA virus belonging to the *Malacoherpesviridae*,
40 poses a significant threat to the global oyster farming industry by initiating a polymicrobial diseases, the
41 Pacific Oyster Mortality Syndrome (POMS, de Lorgeril et al., 2018). In recent years, mortality outbreaks
42 associated with the POMS have caused substantial economic losses. Among several factors (i.e.
43 temperature and food) driving windows of oyster permissivity to diseases oyster age is essential with
44 individuals being susceptible up to ~18 month old (Azéma, Lamy, et al., 2017; Burioli et al., 2017;
45 Nicolas et al., 1992; Renault et al., 1994). Meanwhile, oyster farmers reported mortality associated with
46 OsHV-1 mainly in spat, as survivors demonstrate genetic and epigenetic resistance upon subsequent
47 exposure to the virus (Evans et al., 2017; Gawra et al., 2023).

48 The first comprehensive characterization of the OsHV-1 genome was published in 2005 (Davison et al.,
49 2005). The genome of OsHV-1 spans approximately 207 kilobase pairs (kb) in length and exhibits a
50 distinctive genomic organization comprising unique regions (U_L/U_S : unique long or short) flanked by
51 repeated regions (TR_L/TR_S : terminal repeat long or short and IR_L/IR_S : inverted repeat long or short). The
52 architecture of the OsHV-1 genome can be summarized as $TR_L-U_L-IR_L-X-IR_S-U_S-TR_S$ (Davison et al.,
53 2005). Advances in high-throughput sequencing have facilitated the genomic characterization of several
54 OsHV-1 genotypes, including OsHV-1 μ Var (Burioli et al., 2017), OsHV-1 PT (Abadi et al., 2018),
55 and OsHV-1 SB (Xia et al., 2015). Currently, 28 complete genomes are available in public databases
56 (Delmotte-Pelletier et al., 2022; Morga-Jacquot et al., 2021, Pelletier et al., 2024)). These genomic
57 datasets have revealed a spatiotemporal structuration of viral genotypes in France (Delmotte-Pelletier et
58 al., 2022) and globally (Morga-Jacquot et al., 2021), consistent with oyster farming practices.
59 Furthermore, a recent study has highlighted differentiation between genotypes infecting *M. gigas* and
60 *O. edulis*, with a greater viral diversity observed in *M. gigas* (Pelletier et al., 2024). However, despite
61 significant advancements in understanding OsHV-1 epidemiological and evolutionary processes,
62 several knowledge gaps persist regarding its evolutionary mechanisms and how it adapts to host genetic
63 backgrounds. Indeed, substantial genetic and epigenetic variation for resistance to OsHV-1 has been
64 reported worldwide allowing selective breeding program to sustain oyster production (Camara et al.,

65 2017; Dégremonet et al., 2015b; Divilov et al., 2019; Gutierrez et al., 2020, Gawra et al., 2023; Valdivieso
66 et al., 2025).

67 The evolution of viruses is driven by a range of processes, including mutation, genetic recombination,
68 genetic drift and natural selection. RNA viruses provide vivid examples of high mutation rates and rapid
69 evolution. These viruses are known for their ability to accumulate genetic changes swiftly, resulting in
70 substantial levels of viral polymorphism (Sanjuán, 2012; Sanjuán & Domingo-Calap, 2016, 2016). This
71 genetic diversity empowers them to adapt to newly infected cellular environments and develop strategies
72 to evade vaccines and antiviral drugs (Lauring, 2020). Although large dsDNA viruses exhibit lower
73 mutation rates (*i.e.* between 10^{-03} and 10^{-08} nucleotide substitution/site/year, e.g. 5.9×10^{-08} for the Herpes
74 simplex virus) compared to RNA viruses (*i.e.* between 10^{-02} and 10^{-04} nucleotide substitution/site/year)
75 due to their use of high-fidelity proofreading polymerases (Sanjuán, 2012; Sanjuán et al., 2010; Sanjuán
76 & Domingo-Calap, 2016), their genome stability is increasingly controversial because they sometimes
77 display noteworthy genetic diversity (Sanjuán et al., 2016).

78 Viral evolutionary processes can be investigated through various approaches including sequence
79 analyses, metagenomics, phylogenetics, phylodynamics, and even Experimental Evolution (EE). This
80 former approach involves subjecting viruses to controlled laboratory conditions, enabling the real-time
81 observation and manipulation of evolutionary processes (Kawecki et al., 2012). The emergence and
82 dynamics of mutations, immune escape variants, and evolutionary trade-offs could be assessed by
83 monitoring viral populations under different selective pressures, such as antiviral drug treatments or
84 interactions with immune systems (Kawecki et al., 2012).

85 While most studies using EE focus on easily replicable model organisms such as *Escherichia coli*,
86 *Drosophila spp*, *Saccharomyces cerevisiae* or various bacteriophages (Kawecki et al., 2012), recent
87 research has expanded EE to diverse viruses, including Herpesviruses (Fuandila et al., 2022; Kuny et
88 al., 2020).

89 For instance, *in vitro* EE conducted with populations of Herpes Simplex Virus type 1 (HSV-1) have
90 revealed the accumulation of minor genetic variants and changes in population diversity. Although

91 minor, these changes have been observed to lead to relatively rapid modifications in the viral population
92 at the consensus level (Kuny et al., 2020). Another EE study, conducted on Koi herpesvirus (KHV),
93 which infects the common carp *Cyprinus carpio*, has highlighted the key role of structural variations in
94 the rapid virus evolution *in vitro* (Fuandila et al., 2022). By driving genetic diversity, adaptation, and
95 pathogenicity changes, these large-scale genomic alterations, including deletions, insertions,
96 duplications, and inversions, can rapidly reshape viral genomes, influencing key functional regions
97 (Fuandila et al., 2022).

98 In the absence of oyster cell lines for OsHV-1 propagation, we were forced to use *in vivo* techniques
99 requiring high-throughput sequencing to efficiently and comprehensively analyze the evolution of viral
100 populations. This however offer the advantage to study this host pathogen interaction as a whole in an
101 ecologically relevant experiment.

102 In the context of OsHV-1, our understanding of the evolutionary processes contributing to the diversity
103 of the viral population remains limited. The primary objective of the present study is to gain insights
104 into the evolutionary mechanisms driving the adaptation and diversification of the OsHV-1 genotypes
105 based on infection susceptibility of oysters. To achieve this goal, we used an EE approach spanning up
106 to 28 generations under controlled conditions, combined with a comprehensive sequencing analysis of
107 90 OsHV-1 genomes.

108

109 **Material and methods**

110 **Oysters production**

111 To explore the impact of oyster genetic backgrounds on the evolutionary dynamics of OsHV-1, three
112 oyster populations with varying levels of resistance against OsHV-1 infection were generated. In
113 December 2021, wild oysters were collected from two sites in Charente-Maritime (France) based on
114 their proximity to oyster farms.

115 The first site, La Floride (Marennes-Oléron bay, Charente-Maritime, Lat: 45.803; Long.: -1.153) is
116 located in an area surrounded by oyster farms and densely populated by wild oyster beds. It is annually
117 exposed to OsHV-1 infections (Dégremont et al., 2019). This population is highly resistant to POMS
118 (Valdivieso et al., 2025).

119 The second site, Chaucre (Charente-maritime, Lat: 45.982; Long: -1.396), is located on the West coast
120 of Oléron island in a non-farming area where wild oyster beds are lowly populated. Oysters collected
121 from Chaucre are less likely to be annually infected with OsHV-1 than oysters from densely populated
122 area and may therefore be moderately susceptible to OsHV-1 infection.

123 Moreover, a control oyster population currently used in other studies was included in our experiment
124 (Gawra et al., 2023; Valdivieso et al., 2025). These oysters were initially sampled in 2008 before the
125 emergence of OsHV-1 μ Var, and since 2009, they have been reproduced over six successive generations
126 using oysters from the second generation that have never been exposed to OsHV-1. Therefore, it can be
127 assumed that this oyster population is naive to OsHV-1 since 2009, as animals have always been kept
128 in a secured facility using UV-treated seawater (40 mj/cm²). This population has been described as
129 highly susceptible to OsHV-1, with mortality exceeding 80% (Valdivieso et al., 2025).

130 The three oyster populations (La Floride, Chaucre, and a naïve population) were housed separately at
131 the Ifremer hatchery in La Tremblade (December 2021) to prevent potential pathogen transmission
132 (OsHV-1, *Vibrio aestuarianus*) (Dégremont et al., 2005). Seawater temperature was raised from 11°C
133 to 20°C and maintained under a cultured phytoplankton diet (*Isochrysis galbana*, *Tetraselmis suecica*
134 and *Skeletonema costatum*) to induce gametogenesis (Dégremont et al., 2005).

135 In March 2022, 30 oysters per population were sexed, and gametes were collected via gonad stripping.
136 Female gametes were pooled, filtered (20- μ m and 100- μ m screens), and fertilized with sperm collected
137 individually from males. After 10 minutes, all eggs were combined and transferred to 30 L tanks to
138 minimize sperm competition and maximize effective population size (Boudry et al., 2002). Larval and
139 spat culture followed established protocols (Dégremont et al., 2005, 2007). Progenies were maintained
140 in UV-treated seawater before experimental OsHV-1 infections and renamed FA, NFA, and C, denoting

141 farming area (high density with the typical oyster bed of hundreds individuals/m² and annual POMS
142 reported), non-farming area (low density, less than 20 individuals/m² and no POMS reported), and
143 control populations (laboratory population, highly sensitive to OsHV-1 infection).

144 Before the experiment, spat were acclimated in 120 L tanks with a continuous flow of UV-filtered
145 seawater heated at 19°C. The seawater was enriched with a phytoplankton mixture containing *I.*
146 *galbana*, *T. suecica* and *S. costatum*. The acclimation process spanned at least 2 weeks to ensure optimal
147 growth and feeding conditions, essential for effective virus replication and oyster mortalities (Azéma,
148 Maurouard, et al., 2017).

149 Viral suspension preparation

150 To generate a diversified viral inoculum for infecting all oyster populations, nine viral suspensions were
151 produced between the years 2020 and 2021. These viral suspensions were generated using spat oysters
152 collected during monitoring of OsHV-1 infections from various areas in France. Briefly, approximately
153 1000 specific-pathogen-free (SPF) spat oysters, reared in the Ifremer experimental facilities located in
154 Argenton (Brittany, France), following a standardized methodology (Petton et al., 2015), were deployed
155 into four oyster farming areas during periods of disease outbreaks (Figure 1). These areas included *i*)
156 Brest harbor (BR) (used to produce 2 viral suspensions) (Logonna-Daoulas, lat.: 48.335 long.: -4.318),
157 *ii*) Marennes-Oléron bay (MO) (used to produce 4 viral suspensions) (La Floride, lat.: 45.803 and long.:
158 -1.153), *iii*) Arcachon basin (ARC) (used to produce 2 viral suspensions) (Grahude, lat.: 44.653 and
159 long.: -1.073), *iv*) Leucate lagoon (LEU) (used to produce 1 viral suspensions) (Leucate, lat.: 43.379
160 and long.: 3.571). Seven days after SPF spat were transferred into the four locations, and as mortality
161 began to occur, oysters were brought back to the laboratory and maintained at a temperature of 20°C for
162 7 days. During this period, moribund oysters were systematically sampled daily and then stored at -
163 80°C.

164 In total, nine viral suspensions were prepared according to an established protocol (Schikorski et al.,
165 2011) (Figure 1, Table S1).

166

167



Figure 1: Origin of samples collected to produce viral suspensions.

The samples used to produce the nine viral suspensions were collected between 2020 and 2021 from four locations: Brest harbor (red point), Marennnes-Oléron basin (orange point), Arcachon bay (blue point) and Leucate lagoon (green point).

168

169 Inoculum viral load quantification and viral suspension mix setup.

170 To prepare an equimolar viral inoculum for injection into oysters, viral load of each of the nine viral
171 suspension was assessed. To do so, DNA was extracted from 100 μ l of each viral suspension using the
172 MagAttract[®] HMW DNA kit according to manufacturer's protocol. DNA purity and concentration were
173 assessed by Nano-Drop ND-1000 spectrometer (Thermo Scientific) and Qubit[®] dsDNA BR assay kits
174 (Molecular Probes Life Technologies), respectively. Then quantification of viral copies was carried out
175 by quantitative PCR using a Mx3005 P Thermocycler (Agilent) (Pepin et al. 2008). The outcomes were
176 expressed as the log-transformed copy number of viral DNA per microliter of sea water (cp/ μ L).

177 The nine viral suspensions were pooled to produce the initial viral inoculum (Table S1). To account for
178 variability in the viral copy numbers of each suspension, the pool was generated to ensure equimolarity
179 of the viral copy numbers (Table S1).

180 Experimental evolution design

181 The EE was conducted under controlled laboratory conditions in a Type 2 laboratory at Ifremer's La
182 Tremblade facilities. To initiate the experimental infections, 50 oysters from each oyster populations

183 underwent prior myorelaxation using hexahydrate MgCl_2 (50 g/L) (Suquet et al., 2009). For each oyster
184 population, oysters were individually injected with a 26-gauge needle attached to a multi-dispensing
185 hand pipette. They received 100 μL of pooled inoculum into the abductor muscle (Figure 2, step 1).
186 Simultaneously, for each oyster population, a group of 10 oysters underwent myorelaxation and were
187 injected with artificial sea water, used as a control. The experiment was conducted in duplicate, (*i.e.* 2
188 batches of oysters in 2 tanks) and placed in 5 L plastic tanks filled with seawater at a temperature of
189 20°C and with a phytoplankton mixture (*Isochrysis galbana*, *Tetraselmis suecica* and *Skeletonema*
190 *costatum*). Daily mortality monitoring was carried out for 7 days.

191 After the initial infection, nine tanks were set up: two with infected oysters and one with ASW-injected
192 oysters for each of the three oyster populations. These nine tanks served as the starting point, referred
193 to as "Generation 0" (Figure 2, step 2). To create the first generation of infection (G1) through
194 cohabitation, 25 healthy oysters from each oyster population were introduced 24 hours after infecting
195 the donor oysters (Figure 2, step 3). These healthy oysters were placed in their respective oyster
196 population tank (*i.e.*, 25 individuals from the FA oyster population added to each FA oyster population's
197 infection replicate, and so on). After introducing the healthy oysters to the contaminated tank, they were
198 moved to fresh, pathogen-free seawater 24 hours later to release viral particles (Figure 2, step 4),
199 initiating a second generation of infection. The process of adding healthy oysters and moving them for
200 excretion was repeated for the second generation, following the same procedure as outlined above.
201 During this period, the donor oysters from the cohabitation step were monitored for 7 days to estimate
202 mortality rates (Figure 2, step 5), and all moribund oysters were collected for DNA extraction (see
203 below). In total, we performed 13, 18, and 28 generations for the FA, C, and NFA oyster populations,
204 respectively.

205

206 DNA extraction and sequencing

207 DNA extraction was performed on oysters collected during mortality monitoring (Figure 2, step 5) at
208 generations G0, G5, G10, and G13 for the FA oyster population, at generations G0, G5, G10, G15, and

209 G18 for the C oyster population, and at generations G0, G5, G10, G15, G20, G24, and G28 for the NFA
210 oyster population. For each oyster population, infection generation, and replicate, seven samples of 35
211 mg of mantle tissue were collected for DNA extractions. DNA extractions were performed using the
212 MagAttract® HMW DNA kit according to manufacturer's protocols. DNA purity and concentration
213 were assessed with Nano-Drop ND-1000 spectrometer (Thermo Scientific) and Qubit® dsDNA BR
214 assay kits (Molecular Probes Life Technologies), respectively. Then, quantification of viral copies was
215 carried out by quantitative PCR using a Mx3005 P Thermocycler (Agilent), as detailed earlier. DNA
216 samples from three individuals in each oyster population and replicate were selected based on viral load
217 and DNA concentration. Overall, 96 samples were sequenced by DNA-seq Illumina by IntegraGen SA
218 (Evry, France). PCR free libraries were prepared with the Twist library Preparation Enzymatic
219 Fragmentation (EF) Kit 1.0 (Twist Bioscience) according to supplier recommendations. In brief,
220 following a specific double strand gDNA quantification, 400ng of DNA products were processed. An
221 enzymatic fragmentation was performed to obtain approximately 400bp inserts fragments length. After
222 enzymatic fragmentation, end-repair, A-tailing, ligation to UDI Illumina adapters, libraries were purified
223 and sized with SPRI beads. Sized libraries were then quantified by qPCR and sequenced on the Illumina
224 NovaSeq sequencer as paired-end 150 bp reads.

225

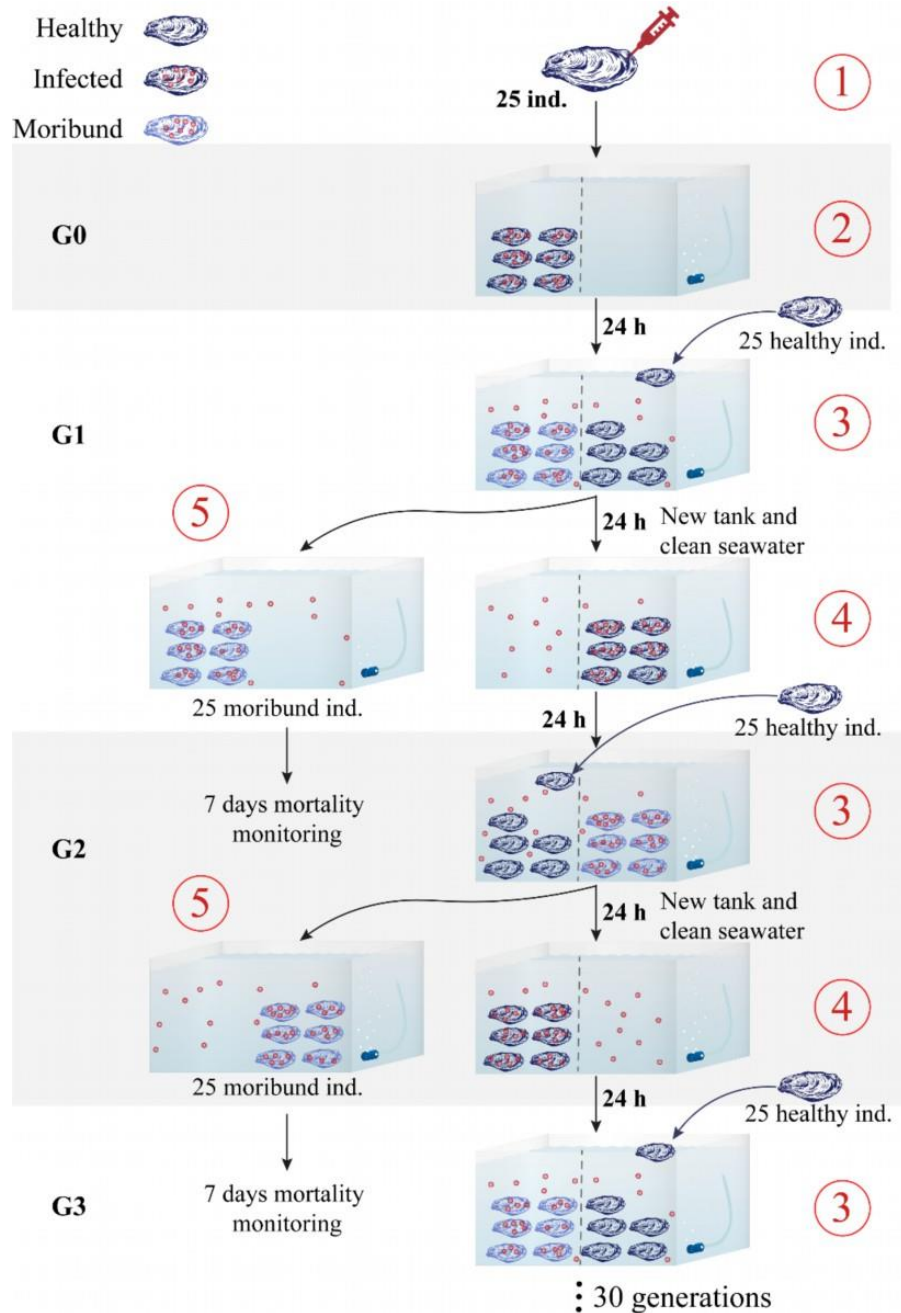


Figure 2: Experimental evolution design for one oyster population.

The experimental evolution was conducted using three oyster populations. Healthy oysters are represented in dark blue, infected oysters in dark blue with red points, and moribund oysters in light blue with red points. All experiments were performed in duplicate. The experimental steps included 1) infection of 25 healthy oysters via intramuscular injection, 2) placement of infected oysters in a tank with clean seawater to initiate viral particle production, 3) addition of 25 new healthy oysters to the tank 24 hours post-infection for exposure, 4) transfer of newly infected oysters to a new tank with clean seawater 24 hours after cohabitation to promote viral particle release, 5) counting of moribund oysters over seven days to estimate mortality rates.

226

227 Assembly of a global consensus for each viral suspension

228 To obtain an overall consensus for each of the viral suspensions, viruses have been sequenced from
229 infected oyster tissue from the same batches used for viral suspension preparation. DNA extractions,
230 purity, concentration and quantification of viral copies were assessed as described previously (see DNA
231 extraction and sequencing). DNA libraries were prepared using a Shotgun PCR-free library preparation
232 kit (Lucigen) and were sequenced using Illumina NovaSeq™ 6000 device (paired-ends, 150 bp) by the
233 Genome Quebec Company (Genome Quebec Innovation Center, McGill University, Montreal, Canada).
234 Raw data have been deposited in the SRA database under Bioproject PRJNA1216400 with accession
235 numbers SAMN46433918 to SAMN46434013 for future reference and accessibility (Table S2).

236 *De novo* OsHV-1 NR-genome assemblies were obtained by processing sequenced reads as previously
237 described (see Dotto-Maurel et al., 2022 for details on the bioinformatic pipeline).

238 To establish a consensus representative of the overall viral suspension used to infect oysters, we
239 performed a multiple alignment of these 126 previously assembled genomes (Figure 3) using MAFFT
240 v1.4.0 (Katoh et al., 2002). The consensus sequence was derived from the alignment and is composed
241 by all the major alleles of OsHV-1 genotypes within the viral suspension. In the forthcoming sections
242 of this manuscript, this consensus will be referred as the “artificial OsHV-1 major NR-consensus”.

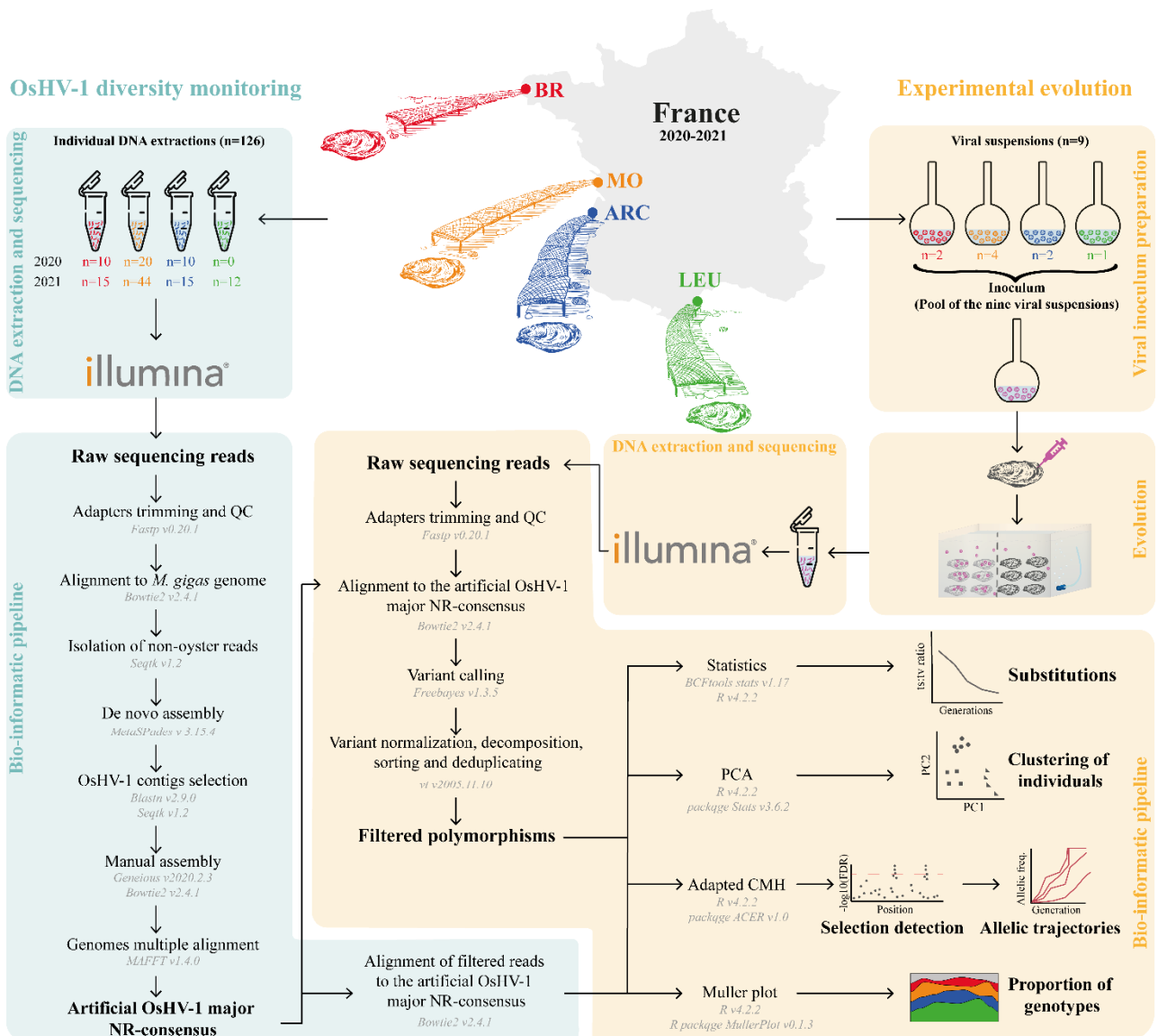


Figure 3: Bioinformatic pipeline used to analyze the dataset and detect selection patterns within OsHV-1 genomes

Steps in blue represent the monitoring of OsHV-1 diversity in France between 2020 and 2021. Pieces of mantle collected from moribund oysters were used to extract DNA, which was sequenced using Illumina technology. Raw reads were assembled *de novo* into OsHV-1 NR-genomes through a specific bioinformatic pipeline previously developed for OsHV-1 genomic analysis (Dotto-Maurel-Pelletier et al., 2019).

Steps in orange represent the analysis of experimental evolution samples and the detection of selection patterns. Moribund oysters collected between 2020 and 2021 were used to produce nine viral suspensions, which were pooled based on equimolar viral copy numbers and named “Inoculum.” Experimental evolution was conducted using this inoculum as the primary source of infection, as described in Figure 2. Pieces of mantle from moribund oysters were used to extract DNA, which was sequenced using Illumina technology. A specific bioinformatic pipeline was developed to detect and characterize traces of selection pressure.

244 Variant calling

245 The variant calling step was performed using an adapted approach previously employed to detect
246 polymorphisms (Figure 3, see Delmotte-Pelletier et al., 2022 for the detailed bioinformatic pipeline).

247

248 Analysis of substitutions types and clustering of individuals based on SNPs

249 To obtain detailed information about transitions (*i.e.* substitution of a purine by a purine or a pyrimidine
250 by a pyrimidine) and transversions (*i.e.* substitution of a purine by a pyrimidine or a pyrimidine by a
251 purine) number within OsHV-1 genome across generations and among host oyster populations,
252 polymorphisms detected in all libraries were used to compute statistics with BCFtools stats v1.17
253 (Figure 3, Danecek et al., 2021). Substitution statistics were averaged by generation and by oyster
254 population and were then plotted using R v4.2.2 and the ggplot package v3.4.0 (Wickham, 2009).

255

256 Exploring sample clustering and SNP data quality assessment through Principal Component 257 Analysis

258 In the context of EE studies, Principal Component Analysis (PCA) is employed to assess sample
259 clustering and evaluate the quality of Single Nucleotide Polymorphisms (SNPs) data, aiding in
260 uncovering underlying population structure and identifying potential data anomalies (Patterson et al.,
261 2006).

262 Utilizing SNP frequencies, the individuals have been projected onto a bidimensional space to check
263 samples clustering. The PCA analysis was performed with the function pcrcomp of the factoextra v1.0.7
264 package (Kassambara & Mundt, 2020) in R v4.2.2 (Figure 3). This information was linked to those of
265 oyster populations, infection generation and replicates.

266

267 Selection detection

268 Selection patterns were detected using a SNP-based method. As the experiment was performed in
269 duplicates for three oyster populations, selected SNPs could be detected within each oyster population
270 in all replicates or in specific replicates. Selection of particular OsHV-1 genotypes within the different
271 oyster populations is independent, and detected SNPs were not comparable among oyster populations.
272 Consequently, analyses were carried out for each oyster population independently. Initially, to identify
273 SNPs demonstrating significant shifts in allele frequency (AF), a comparison was made between the
274 initial (G0) and evolved populations (reaching G13, G18, and G28 for FA, C and NFA oyster
275 populations, respectively). To achieve this, an adapted Cochran-Mantel-Haenszel (CMH) test was used
276 (Figure 3), which calculated p-values for each SNP detected within the datasets using the ACER v1.0
277 package (Barghi et al., 2020; Spitzer et al., 2020) in R. These p-values reflect the variance in SNP
278 frequencies across generations while accounting for the influences of genetic drift and sampling (Spitzer
279 et al., 2020). Subsequent to this, the p-values underwent adjustment using the false discovery rate (FDR)
280 method, as implemented in the p.adjust function from the stats v4.2.2 package (R Development Core
281 Team, 2005). The resultant adjusted p-values were then plotted onto the OsHV-1 genome using a
282 Manhattan plot, using the ggplot2 v3.4.0 package (Wickham, 2009). Alleles were considered as under
283 selection when p-values were higher than the CMH cutoff determined as 5% of false discovery rate.
284 Finally, using the adjusted p-values and the allelic frequencies of SNPs across generations, allelic
285 frequency trajectories were generated for each replicate. This visualization was achieved using the
286 ggplot v3.4.0 package (Wickham, 2009).

287 Genotypes proportions

288 A Muller plot was used to visualize the emergence or loss of genotypes specific to the origin of the
289 initial viral suspension used to infect the oysters. As described previously, the initial viral suspension
290 injected into oysters at G0 was produced by pooling of nine viral suspensions. Libraries associated with
291 the sequencing of the individuals collected during the same mortality events within each basin were used
292 to determined SNPs specific to each origin. Raw reads were then filtered and trimmed as described

293 previously, and aligned to the “artificial OsHV-1 major NR-consensus”. Variant were called using
294 Freebayes and named using unique identifier as described above (Figure 3).

295 Each sample was collected from a specific location, allowing each SNP detected in the libraries to be
296 assigned accordingly. SNPs identified during the EE of OsHV-1 were classified by their origin and used
297 to visualize their relative abundance within each origin using the MullerPlot package v0.13 (Farahpour
298 et al., 2022) in R (Figure 3).

299

300 Results

301 Evolution of survival rates and viral load in Pacific Oyster spat infected with OsHV-1 across 302 generations

303 To assess the impact of the infection by the pool of OsHV-1 genotypes on Pacific oyster spat, the
304 survival rate was monitored for 7 days post-infection (dpi) in both infected replicates and control tanks
305 for each of the three oyster populations. No oyster mortality was observed in the conditions without
306 viral infection (*i.e.* ASW-injected oysters) throughout the entire experiment.

307 Cohabitation-induced infection relies on viral propagation within the host. Challenges in sustaining
308 reinfection through successive generations led to the termination of experimental evolution at the 13th
309 generation for the farming area (FA) population, the 18th for the control (C) population, and the 28th for
310 the non-farming area (NFA) population.

311 Mortality rates fluctuated across generations but showed an overall decline over the course of
312 experiment (Figure 4A). At G0, mortality rates were 88%, 96%, and 84% respectively for FA, C and
313 NFA oyster populations. In the C population, mortality steadily decreased to 42% at G15 before rising
314 to 56% at G18, at which point the experiment ended due to the absence of mortality at G19.

315 In contrast, FA and NFA populations showed variable mortality trends. FA oysters experienced a sharp
316 drop of mortalities to 58% at G5, a rise to 87% at G10, and a drastic decline to 2% thereafter, leading to

317 experiment termination. NFA oysters followed a similar pattern beginning with a decrease of mortalities
318 to 50% at G5, a spike to 99% mortality at G10, then fluctuations between 28% and 89% mortality.

319 Variations in viral loads were observed both among individuals and among oyster populations (Figure
320 4B). The mean viral load within individuals from the oyster populations FA exhibited a slight increase
321 across generations, rising from $8.38 \times 10^5 \pm 5.49 \times 10^5$ viral DNA copies/ng of DNA at G0 to $1.13 \times 10^6 \pm$
322 4.81×10^5 viral DNA copies/ng of DNA (Figure 4B). Conversely, the average viral load within
323 individuals from the oyster populations C and NFA demonstrated a slight decrease across generations.
324 For the population C, the viral load varied from $6.62 \times 10^5 \pm 4.49 \times 10^5$ viral DNA copies/ng of DNA at
325 G0 to $3.34 \times 10^5 \pm 2.54 \times 10^5$ viral DNA copies/ng of DNA at G18. In the case of the population NFA, the
326 viral load ranged from $2.09 \times 10^6 \pm 1.20 \times 10^6$ viral DNA copies/ng of DNA at G0 to $9.35 \times 10^5 \pm 1.11 \times 10^6$
327 viral DNA copies/ng of DNA at G28 (Figure 4B).

328

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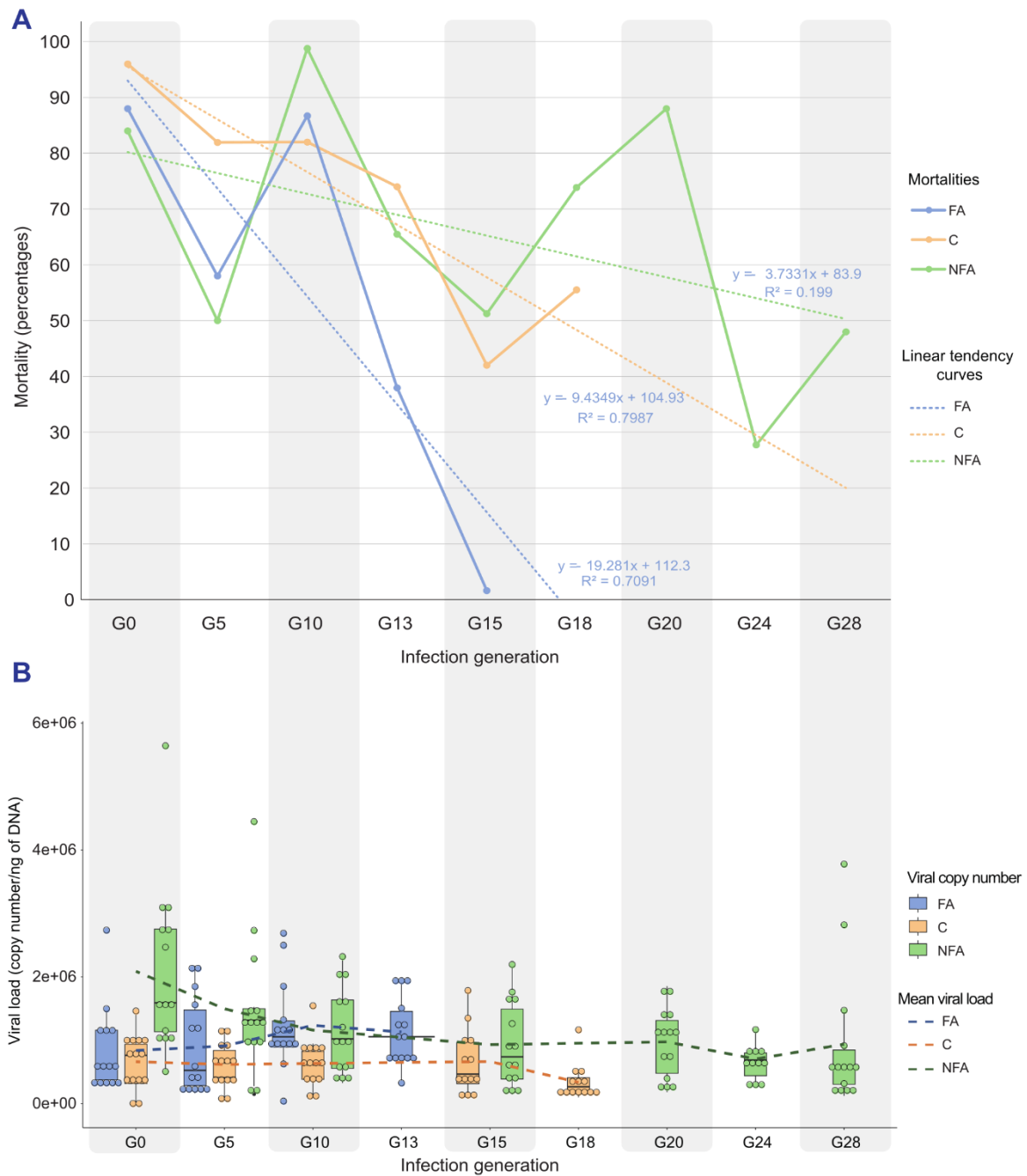


Figure 4: Oyster mortalities and viral load from the three oyster populations at generations 0, 5, 10, 13, 15, 18, 20, 24 and 28.

A) Oyster mortality was monitored daily to estimate mortality rates within each oyster population for selected generations. The solid curves and dotted lines correspond to mortality rates and the linear tendency of mortality rates respectively, for oysters from the line FA oyster population (in blue), from the C oyster population (in orange), and from the NFA oyster population (in green).

B) Viral loads were assessed from seven individuals for each generation, oyster population, and replicate to monitor the evolution of the infection over generations. Viral loads of individuals are represented by points (in blue, orange, and green for oysters from lines FA, C, and NFA, respectively). The box plots represent median values, the first and third quartiles, and the standard deviation of viral loads within an oyster population for a specific generation. The dotted lines illustrate the viral load for oysters from the FA oyster population (in blue), from the C oyster population (in orange), and from the NFA oyster population (in green).

331 Dynamics of substitutions and transition / transversion ratio

332 Studying substitution patterns across viral generations in EE is crucial for understanding evolutionary
333 dynamics, genetic diversity, and adaptation (Bromham, 2020).

334 A prominent trend emerges for all three oyster populations, indicating a rapid reduction in the count of
335 SNPs in the OsHV-1 genome across the populations throughout the generations of infection (Figure 5).
336 The trend is more pronounced for the NFA oyster population extending from 372 ± 55 SNPs at G0 to
337 294 ± 43 SNPs at G5 (Figure 5A).

338 A statistical analysis, based on the identified SNPs, was conducted to assess the transition / transversion
339 (ts/tv) bias (Figure 5B). The ts/tv ratio of the virus remained consistently below 1 across all generations
340 and lines, except for the samples collected from the FA oyster populations at G0, for which the virus
341 displayed a ts/tv ratio of 1.26, indicating an excess of transitions (Figure 5B). Additionally, the viral
342 ts/tv ratio demonstrates a decreasing trend over generations for both the FA and C oyster populations.
343 Regarding the NFA oyster populations, the overarching pattern leans towards a decline in the ts/tv ratio;
344 however, an upward shift in the ratio is noticeable in G10.

345

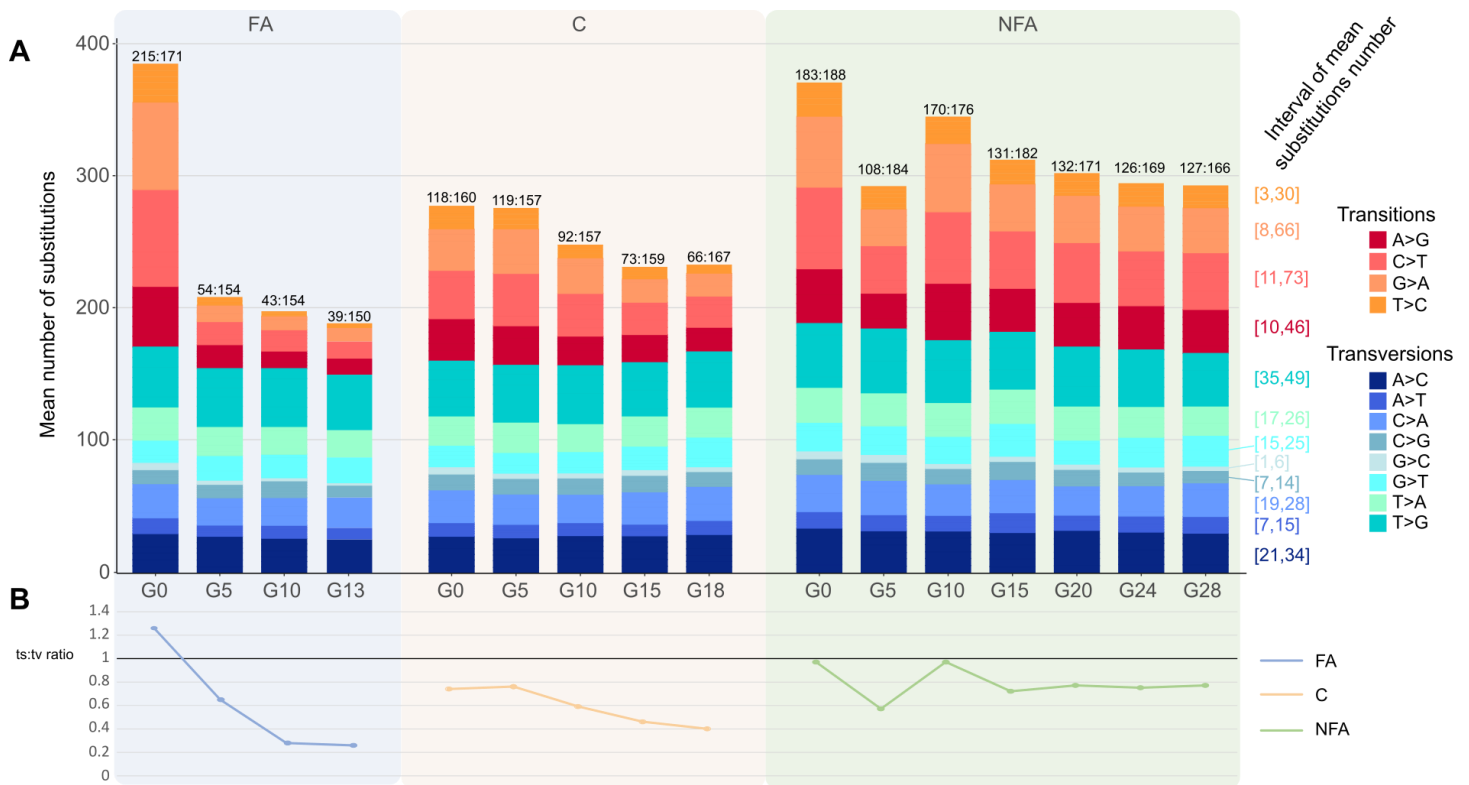


Figure 5 : Substitution types and transition:transversion ratio observed within *OsHV-1* populations.

A) The mean number of viral substitution types per generation and oyster population is shown. Transitions (substitutions of a pyrimidine by a pyrimidine or a purine by a purine) are represented by a gradient from red to orange, while transversions (substitutions of a purine by a pyrimidine or vice versa) are shown with a gradient from blue to green. The numbers above the stacked bars indicate the counts of transitions and transversions, while the numbers in square brackets to the right of the bars represent the average number of each substitution type in the samples.

B) Curves illustrating variations in the ts:tv ratio across generations for each oyster population. Line colors correspond to the different oyster populations.

346 Unveiling *OsHV-1* population structure and anomalies in experimental evolution through SNP

347 Data analysis and principal component projection

348 Using SNP frequencies, we projected individuals *OsHV-1* genomes onto a bidimensional space (Figure

349 6). The first principal component axis accounted for 12.61% of the variance in the data, while the second

350 principal component axis explains 9.89% of the variance (Figure 6). The first principal component aligns

351 with the progression of infection generations, while the second component is indicative of oyster

352 population associations or susceptibility traits within oyster populations. The graphical representation

353 reveals the presence of four distinct clusters corresponding to the interplay between oyster populations

354 and infection generations.

355 The first cluster encompasses samples from all oyster populations at G0, with those collected from the
356 replicate 1 of the C population at G0, G5, G10, G15, and G18 (Figure 6A, group 1) suggesting a
357 maintenance of the initial diversity along the generation 0 to 18 in the replicate 1 of the C population.

358 The second cluster (Figure 6A, group 2) encompasses samples primarily collected from FA line at G5,
359 G10, and G13, as well as the replicate 2 from C line at G5, G10, G15, and G18.

360 Finally, the third cluster comprises specimens sourced from the NFA oyster population, further divided
361 into two subsets. The initial subset (Figure 6A, group 3.1) exclusively represents samples obtained
362 during G5, while the second subset (Figure 6A, group 3.2) corresponds to samples collected between
363 G15 and G28.

364

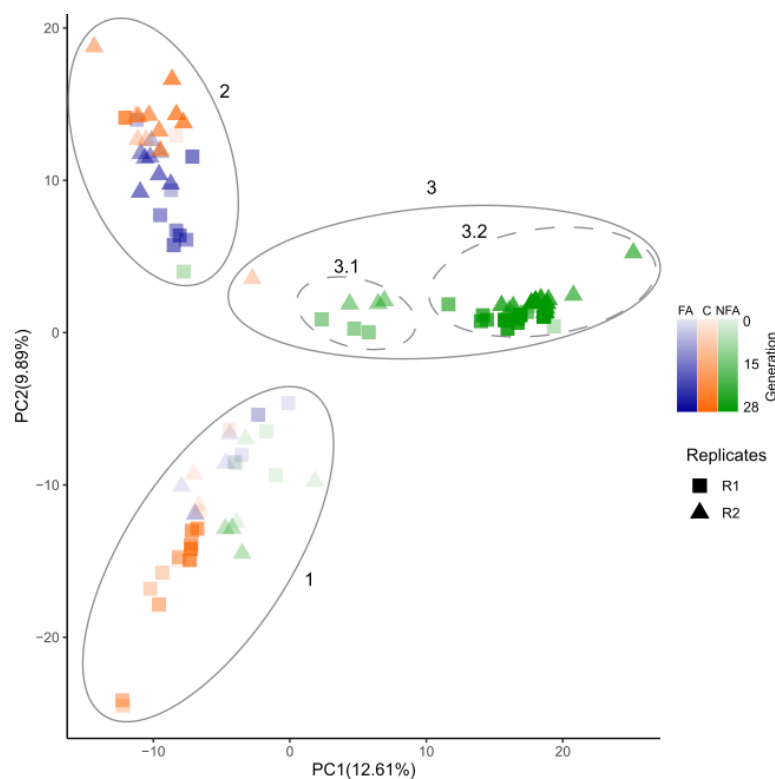


Figure 6 : Principal component analysis reveals *OsHV-1* population structure and generation-oyster populations associations.

Projection of individual similarities onto the first and second principal components. The colors of the points correspond to the infected oyster populations: individuals from the FA oyster population are colored in blue, individuals from the C oyster population are colored in orange, and individuals from the NFA oyster population are colored in green. Infection generations are represented by a gradient of colors, and replicates are indicated by different point types (square for replicate 1 and triangle for replicate 2). The clustering of samples is indicated by ellipses and numbered from 1 to 3. For the cluster 3, two subclusters were identified numbered 3.1 and 3.2.

365

366 Detection of regions under selection pressure and allelic frequency trajectories

367 The Cochran-Mantel-Haenszel test (CMH) is widely used to detect selection pressures by analyzing
368 allele frequency shifts in experimental evolution and time series data (Spitzer et al., 2020).

369 The CMH test, employed on SNPs detected within each viral population collected from the three oyster
370 populations independently (Figure 7), revealed that in oyster populations FA (Figure 7A) and C (Figure
371 7B), no regions were significantly impacted by selection. However, within the NFA oyster populations,
372 a total of 117 positions demonstrated candidate selection signatures (Figure 7C). These positions are
373 distributed across the entire OsHV-1 genome, with a distinct emphasis on specific regions, such as
374 repeated regions or the UL region spanning 50 kb to 80 kb. More precisely, 28 potentially selected
375 positions were located within intergenic regions, including one SNP in the stem loop, and 89 SNPs were
376 located within ORFs (Figure 8, Table S3).

377 Specifically, among the 89 SNPs situated within ORFs, 58 correspond to ORFs with undefined
378 functions, while 12 are found within ORFs encoding transmembrane proteins (Table S3). Additionally,
379 8 SNPs are situated within ORFs encoding secreted proteins, four within ORF 100 coding for a DNA
380 polymerase, three within ORFs encoding Zinc-finger, Ring-type proteins, two within ORFs encoding
381 BIR repeat motifs, one within ORF115 associated with an Origin-binding protein, and the final SNP
382 within ORF20 encoding a ribonucleotide reductase (Table S3).

383

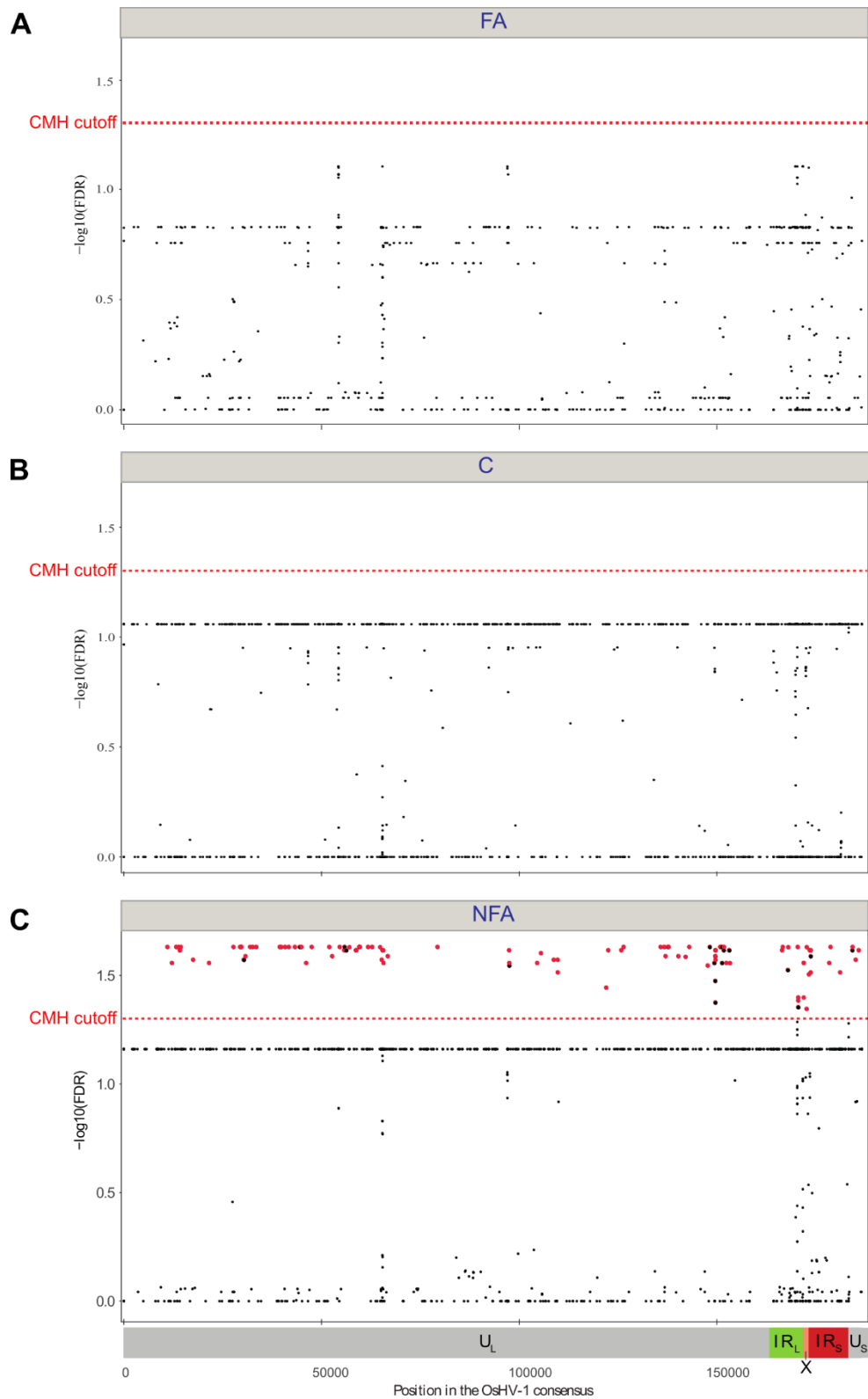


Figure 7: Genomic distribution of candidate SNPs in oyster populations.

The scatter plots show the negative log₁₀-transformed p-values of SNPs corresponding to genomic positions for A) Farming area, B) Control and C) Non-Farming area oyster populations. These p-values were calculated using the CMH test, which involved comparing the viral populations of the founder and evolved states. Significance thresholds are indicated by red dotted lines, representing the CMH cutoff at a 5% false positive rate. Significant p-values are highlighted by red points. The genomic architecture of the OsHV-1 genome is represented below the graphs, depicting unique and repeated regions as follows: U_L/U_S (unique long/short) and IR_L/IR_S (inverted repeat long/short).

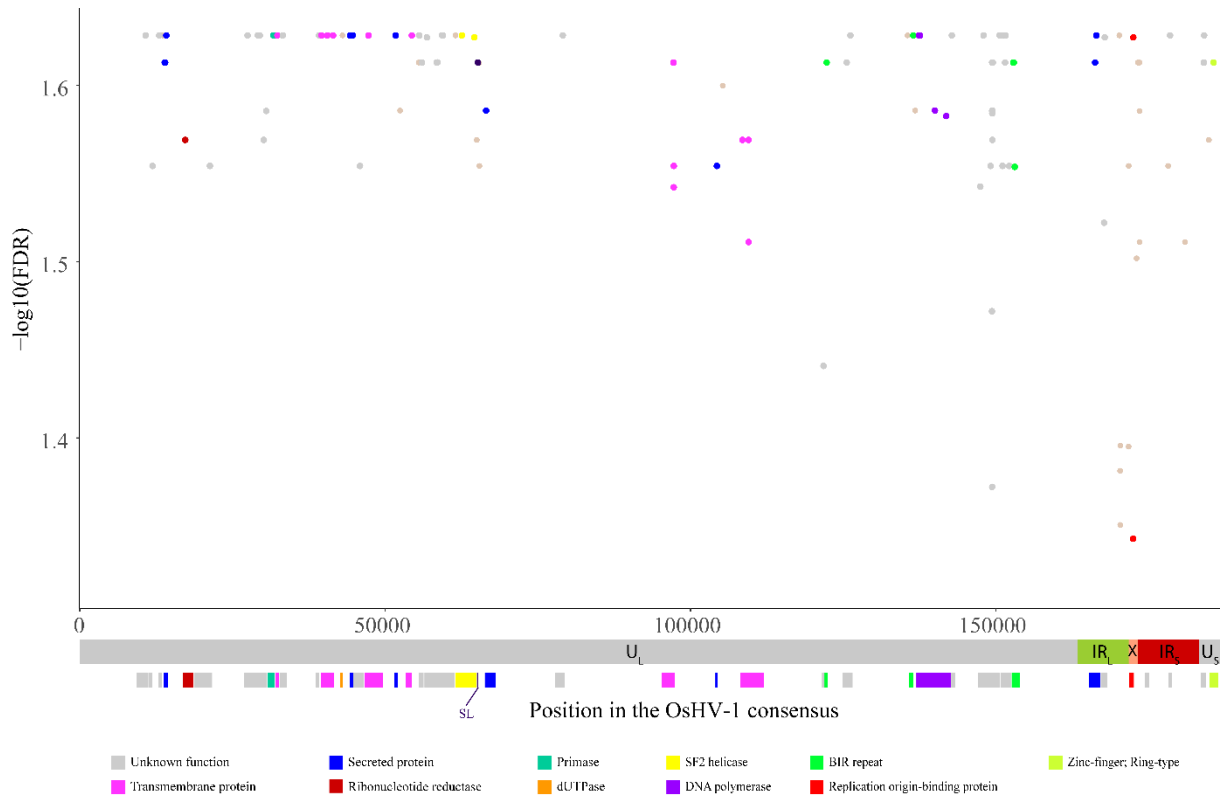


Figure 8: Zoom on the genomic distribution of selected SNPs in Non-Farming area oyster population.

The scatter plot shows the negative log₁₀-transformed p-values of selected SNPs corresponding to genomic positions. These p-values were calculated using the CMH test, which involved comparing the viral populations of the founder and evolved states. Significance thresholds representing the CMH cutoff at a 5% false positive rate are displayed. Below the plots, the genomic structure of the OsHV-1 genome is presented, highlighting distinct segments such as unique and repeated regions denoted as U_L/U_S (unique long/short) and IR_L/IR_S (inverted repeat long/short). Moreover, the genomic architecture is supplemented by the depiction of ORFs (Open Reading Frames) housing the selected SNPs, color-coded based on their encoded functions or domains. SL: Stem-loop. The SNPs themselves are also color-coded according to the specific impact they have on the function of the corresponding ORF.

385

386 An allelic frequency trajectories (AFT) plot was employed for candidate SNPs detected within the NFA
 387 oyster population to track changes in allelic frequencies over generations and get insights into the
 388 evolutionary dynamics within populations, especially for positive selection (Barghi et al., 2020). The
 389 analysis of allelic frequency trajectories has been performed separately for both replicates (Figure 9).

390 The AFT plot of candidate SNPs reveals a general trend of increasing allelic frequencies, ultimately
 391 resulting in the fixation of selected alleles in both replicates. However, in the first replicate, the allelic
 392 frequencies of candidate SNPs increase between G0 and G5, then decrease between G5 and G10, and
 393 finally increase again between G10 and G15 to reach fixation (Figure 9A). In the second replicate, a
 394 similar trend is observed, but only for 77 SNPs (Figure 9B). Moreover, alleles of 66 SNPs for replicate

395 1 and 26 SNPs for replicate 2, represented at the bottom of the plot, show an increase in frequencies,
396 albeit to a lesser extent (Figure 9A and B).

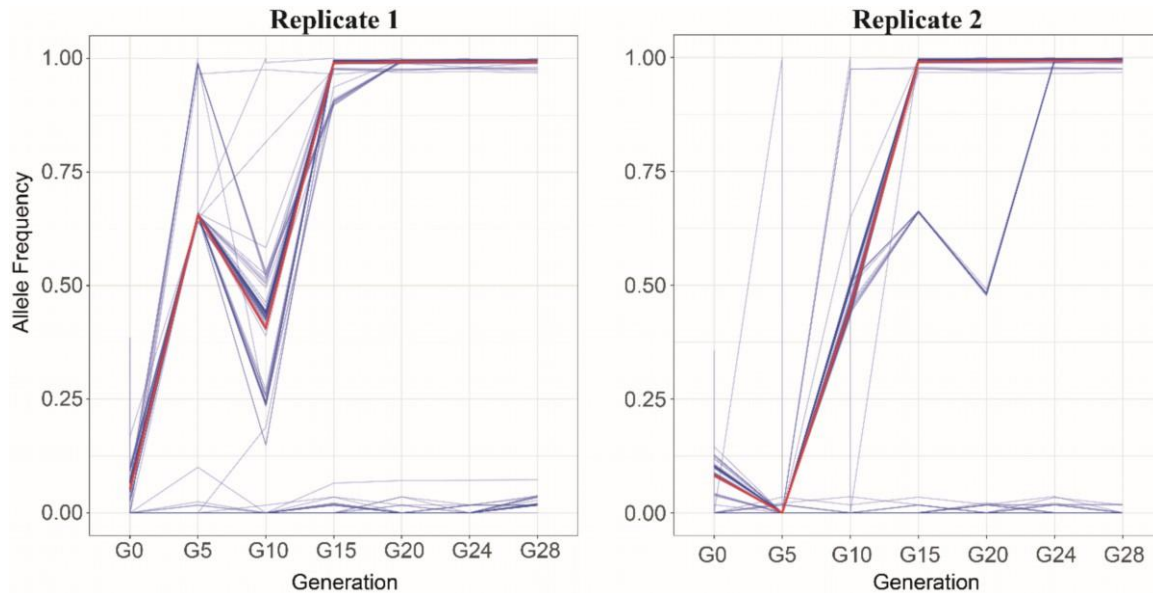


Figure 9: Allelic frequency trajectories plot for the top 100 candidate SNPs detected within the *OsHV-1* samples of replicate 1 and 2 collected from oysters of the NFA oyster population.

Each blue line represents the allelic frequency trajectory of a specific SNP across generations. Red line represents the median allelic frequency trajectory.

397

398 Relative abundance of SNP from specific origin

399 To track the presence of viral genotypes specific to one of the geographic origins and their fate across
400 generations of infection, a Muller plot was generated (Figure 10). The results revealed that multiple
401 genotypes from different origins coexist from the beginning to the end of the experiment within each
402 oyster populations. At the generation 0, the relative abundance (RA) of genotypes specific to MO and
403 LEU is higher than that of genotypes specific to ARC and BR.

404 This trend is generally maintained across generations of infections for all oyster populations. However,
405 depending on the oyster populations, the RA varies over the generations. Specifically, for the FA oyster
406 population, *OsHV-1* SNPs specific to MO are present at a RA of 0.2 in G0, then their RA increases to
407 reach 40% in G10 before slightly decreasing. The trend is generally similar for *OsHV-1* SNPs specific

408 to ARC and BR, but with much lower RA (between 0.02 and 2.0). As for OsHV-1 SNPs specifically
409 from LEU, the trend is reversed. Indeed, the initial RA decreased from 0.4, to 0,1 at G5.

410 Moreover, over the course of the experiment, SNPs that were not initially present in any of the viral
411 suspensions began to emerge. At G0, the RA of these SNPs was relatively low within farming and non-
412 farming oyster populations (RA = 0.1). However, as the infection generations progressed, their RA
413 increased. In contrast, the RA of “other” SNPs within the control oyster populations at the beginning of
414 the experiment was 0.3, and then it experienced fluctuations across generations. First, it decreased to
415 RA=0.2 until G5, then increased to RA=0.5 until G10 and decreased to RA=0.2 until G15.

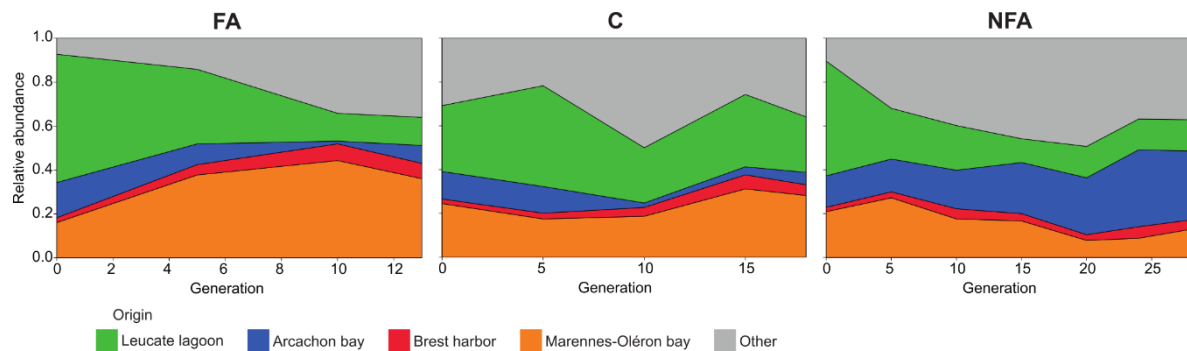


Figure 10 : Muller plot representing the relative abundance of viral SNPs collected from specific origins for each oyster population during the experimental evolution.

The relative abundance of viral SNPs is color-coded according to their geographical origin of the viral suspension: the orange area corresponds to genotypes collected specifically in MO, the red area corresponds to genotypes collected specifically in BR, the blue area corresponds to genotypes collected specifically in ARC, the green area corresponds to genotypes collected specifically in LEU, and the grey area corresponds to genotypes that appeared during the experimental evolution.

416

417

418 Discussion

419 To mitigate the impact of infectious diseases, it is essential to unravel the mechanisms that shape the
420 evolution and dissemination of disease-causing microorganisms (Gu et al., 2021; NIH, 2007).

421 Variability in host resistance levels is known to play a significant role in driving these processes (Gandon
422 & Michalakis, 2000; Kubinak & Potts, 2013). This study provides valuable insights into the intricate
423 evolutionary dynamics of OsHV-1 and its interactions with Pacific oysters, focusing on how the virus
424 adapts and diversifies in response to variations in host genetic backgrounds. These findings shed light

425 into mechanisms driving viral evolution, a crucial step in understanding and addressing the devastating
426 effects of this virus inducing POMS on the oyster farming industry.

427 During this experimental evolution, we observed a decrease in both oyster mortalities and viral load
428 across successive generations of OsHV-1 infection. This decline in mortality rates can be attributed to
429 several factors.

430 First and foremost, it can be explained by the selection of less virulent viral variants. As is well-
431 established, the interactions between a pathogen and its host are characterized by a delicate balance
432 between virulence and transmission (Anderson & May, 1982). In conditions where the oyster population
433 is limited, such as in our experimental setup with 25 individuals per tank, infected hosts succumb
434 rapidly. A plausible strategy adopted by the virus may be to reduce its virulence in order to maintain a
435 stable host population while ensuring efficient replication without lethality. It is a well-documented
436 phenomenon that there is a trade-off between virulence and transmission, where increased transmission
437 is associated with decreased virulence, and conversely, increased virulence leads to decreased
438 transmission (Anderson & May, 1982; Gandon & Michalakis, 2000; Kun et al., 2023). Similar results
439 were obtained in an EE study involving tobacco and pepper plants infected with the tobacco etch
440 potyvirus (TEV). In this study, viral load and virulence significantly decreased across generations, a
441 phenomenon linked to the host lineage, which plays a pivotal role in shaping the fate of viruses (Cuevas
442 et al., 2015).

443 Secondly, the reduction in mortality rates can be attributed to a lack of complementation between
444 genotypes (Froissart et al., 2004; Montville et al., 2005). In fact, host infection rarely occurs with a
445 single genotype, but often involves co-infection, with multiple viral genotypes contributing to the
446 infection and associated symptoms. It has already been shown that certain genotypes that have
447 developed disadvantageous mutations in important genes, can be complemented by other genotypes that
448 do not carry these mutations and whose corresponding proteins remain functional (Froissart et al., 2004;
449 Montville et al., 2005). This mechanism allows low-fitness genotypes to gain a phenotypic advantage
450 by utilizing intracellular proteins produced by co-infecting high-fitness strains (Froissart et al., 2004;
451 Montville et al., 2005). In the context of our study, where genetic diversity has likely been drastically

452 reduced, possibly due to a bottleneck effect, it is possible that some genotypes have been purged from
453 the viral population, leading to limited complementation events and consequently a reduction in
454 virulence over successive generations. Bottlenecks, which occur when a population undergoes a sharp
455 reduction in size due to factors such as transmission events or selective sweeps, can drastically reduce
456 genetic diversity by eliminating rare variants and fixing certain alleles through genetic drift (Zwart &
457 Elena, 2015). In viral populations, this can be particularly impactful, as beneficial interactions between
458 co-infecting genotypes rely on the presence of genetic diversity (Zwart & Elena, 2015). If only a small
459 subset of the original population survives a bottleneck, previously co-existing genotypes that enabled
460 complementation may be lost, preventing the recovery of deleterious mutations and diminishing overall
461 viral fitness (Zwart & Elena, 2015). As a result, the ability of the virus to sustain high virulence through
462 cooperative interactions is compromised, leading to an overall reduction in mortality rates.

463 A third hypothesis to explain declines in mortality rates and viral loads is the temporal shift in the peak
464 of viral particle release into seawater as infection generations progress. Consequently, the viral load
465 would be high when cohabitation begins and progressively decreasing over time. In the case of OsHV-
466 1, the excretion peak by oysters occurs between 24 and 48 hours post-infection (hpi) (Delmotte et al.,
467 2020), but this timing can vary depending on several factors, including oyster susceptibility, the initial
468 viral load in infected oysters, the number of donors, food availability, oyster age, water temperature, and
469 salinity (Dégremont, 2011; Delmotte et al., 2020; Evans et al., 2017; Pernet et al., 2018, 2018; Petton et
470 al., 2015; Schikorski et al., 2011). In our experiment, oysters used for successive infections came from
471 the same cohort, meaning they aged between the start and end of the experimental evolution, potentially
472 altering their susceptibility to infection and the timing of peak viral excretion. Additionally, previous
473 studies have shown that oyster infections via cohabitation lead to fewer fatalities due to a progressive
474 decline in the initial viral load from one infection cycle to the next (Cain et al., 2021). While it may not
475 be the optimal method for EE, successive infection by cohabitation was chosen to mimic the natural
476 infection of oysters by OsHV-1 in the field.

477 Genetic mutations play a pivotal role in shaping the evolutionary landscape of organisms, with
478 transitions and transversions representing two distinct classes of point mutations that contribute to

479 genetic diversity and adaptation. Molecular evolutionary hypothesis suggests that natural selection
480 promotes amino acid substitutions mostly through transitions (Stoltzfus & Norris, 2016). The substantial
481 decline in the count of single nucleotide polymorphisms (SNPs) and the decrease in the
482 transition/transversion (ts/tv) ratio across all three oyster populations over successive infection
483 generations suggest a trend toward genetic homogenization within the viral populations. This trend could
484 arise from selection favoring specific viral genotypes or from a reduction in genetic diversity due to
485 population bottlenecks (McCrone & Luring, 2018). Interestingly, similar patterns of reduction in viral
486 variants have been observed in two distinct EE studies. Firstly, in the case of the Cucumber mosaic virus
487 injected into tobacco plants during an experimental infection, viral populations underwent a population
488 bottleneck due to genetic drift, resulting in a significant reduction in viral diversity (Li & Roossinck,
489 2004). Similarly, when Potato virus Y passed through three different potato plant genotypes with varying
490 levels of susceptibility to the virus, a combination of strong selective sweeps and bottlenecks led to a
491 reduced viral diversity by the fifth generation (Kutnjak et al., 2017). Given the context of OsHV-1
492 infecting different oyster genetic backgrounds and considering host population size, it is plausible that
493 the viral population underwent a bottleneck, leading to a reduction in the number of viral variants.

494 In contrast to the experimental developments carried out on herpesviruses, no structural variation was
495 observed in the genomes of OsHV-1 (Fuandila et al., 2022). The absence of structural variations during
496 short-term experimental evolution studies of viruses can be attributed to the limited timeframe, which
497 may be insufficient for such large-scale genomic changes to emerge or become detectable. Structural
498 variations, including large-scale rearrangements or duplications, often require extended periods to
499 develop and reach appreciable frequencies within viral populations. For instance, studies on RNA
500 viruses have demonstrated that while point mutations can accumulate rapidly, more substantial genomic
501 alterations necessitate longer evolutionary timescales (Duffy et al., 2008; Elena & Lenski, 1997).

502 The PCA analysis of SNP frequencies unveiled distinct clusters that corresponded to the different oyster
503 populations and infection generations, signifying genetic differentiation. Through the process of
504 experimental evolution, viral populations exhibited adaptability across successive generations of
505 infection, with individuals organizing into clusters according to their infection histories. Particularly

506 noteworthy, the clustering of individuals from the non-farming (NFA) and farming area (FA) oyster
507 populations is clearly delineated and distinct, indicating a contrasting evolution within these two
508 populations. In contrast, control oyster population (C) exhibited mixed clustering, likely owing to their
509 intermediate genetic status. These findings strongly suggest ongoing genetic differentiation, which could
510 be attributed to varying levels of oysters susceptibility to OsHV-1 infection (Kubinak & Potts, 2013).
511 Within the NFA oyster population, distinct clusters emerged, indicating evolutionary changes across
512 infection generations in response to selective pressures.

513 The identification of 117 candidate selection signatures in the OsHV-1 genomes, in conjunction with
514 the allelic frequency trajectories (AFT) plots collected from the NFA oyster population, provides
515 compelling evidence of selective pressures acting on OsHV-1 during the course of the experimental
516 evolution. The trend toward increasing allelic frequencies, ultimately leading to allele fixation in both
517 replicates, strongly indicates positive selection (Barghi et al., 2020). Notably, selective pressure was
518 exclusively detected within the OsHV-1 genome during successive infection in the NFA oyster
519 population.

520 One plausible hypothesis to explain this observation is that in more susceptible oysters, viral replication
521 is favored (Dégremont, 2011), leading to an increase in mutation rates and enabling the fixation of
522 advantageous alleles within populations (Elena & Sanjuán, 2005; Peck & Luring, 2018). Given that
523 viral diversity and natural selection processes in viruses are primarily influenced by environmental
524 changes or by the host's immune system acting as a filter to limit viral diversity and adaptation, one
525 might have expected to detect selection signals in viral genomes infecting the FA oyster population
526 (Kubinak & Potts, 2013). However, since no selection signals were detected for the FA oyster
527 population, despite the observed reduction in viral diversity, this strengthens the hypothesis that the viral
528 populations infecting the FA oyster population may have been influenced by a bottleneck effect and
529 genetic drift.

530 These selection signatures are distributed across the viral genome, with a specific emphasis on repetitive
531 segments and the U_L region. The presence of selection signatures within ORFs encoding various proteins
532 suggests that both regulatory and functional elements of the virus are under selective pressure.

533 Specifically, regions responsible for vital biological functions or domains related to the attachment of
534 viral particles to host cells, DNA replication, synthesis, packaging, as well as transmembrane and
535 membrane proteins, could be crucial for the adaptation to resistant or susceptible hosts. It is worth noting
536 that viral entry proteins play a pivotal role in enhancing viral fitness, and even a single amino acid
537 mutation within these proteins can modify the virus's ability to infect hosts with varying levels of
538 susceptibility to infection (van Sluijs et al., 2017). Moreover, studies have revealed that various domains,
539 including envelope domains (such as membrane glycoproteins and transmembrane receptors), auxiliary
540 domains (like Zinc-finger, RING type, dinucleoside kinase), and modulation and control domains (such
541 as Interleukin, Interferon-regulatory factor, or Zinc-finger), have been acquired, duplicated, or lost
542 during the evolution of Herpesviridae (Brito & Pinney, 2020). Many of these acquired domains enabled
543 viruses to specifically bind to host cells and to evade or manipulate the host's immune system. In the
544 present study, the majority of the signals of selection were identified within these domains, indicating a
545 potential adaptation of essential proteins that leads to a specialization of viral genotypes to their
546 respective host genetic background. Additionally, two previous studies conducted on OsHV-1 (Delmotte
547 et al., 2020; Pelletier et al., 2023) have demonstrated that selective pressure plays a key role in the
548 adaptation of the virus to different host species and environments through mutations, particularly in
549 ORFs encoding viral membrane-related and metabolic-related proteins, further supporting the
550 hypothesis of immune selection of OsHV-1 within the NFA oyster population. Based on the results
551 obtained in this study, it is evident that adaptation can also occur at a finer scale within the host's genetic
552 background, which is influenced in this case by the geographic origin of the parental individuals (*i.e.*
553 farmed or not farmed area).

554 Conclusion

555 This study provides valuable insights into the evolutionary forcing mechanisms of OsHV-1 and its
556 interactions with Pacific oysters, with a particular emphasis on viral adaptation and diversification in
557 response to host infection susceptibility. Employing an EE approach, it sheds light on the dynamic nature
558 of host-virus interactions, the potential for viral adaptation, and the potential role of genetic diversity in
559 shaping the outcome of these interactions. The nucleotide evolution of OsHV-1, particularly a decrease

560 in transition and transversion numbers, appears to be mainly driven by both genetic drift and positive
561 selection linked to the oyster genetic background, resulting in a reduction in viral diversity and the
562 fixation of specific alleles within ORFs dedicated to host-virus interactions and virus functional
563 maintenance. Further research into the functional consequences of these changes is necessary. These
564 findings contribute to our understanding of the mechanisms governing viral evolution and hold
565 significant implications for mitigating the devastating impacts of OsHV-1 on oyster farming industries.
566 Future studies should continue to explore the intricate dynamics of host-virus interactions and their
567 implications for viral evolution and disease management.

568

569 **Data availability**

570 Raw data have been deposited on the SRA database under Bioproject PRJNA1216400 accession
571 numbers SAMN46433918 to SAMN46434013 for future reference and accessibility (Table S2). All the
572 scripts are available at https://gitlab.ifremer.fr/lgpmm/experimental_evolution.git.

573

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593 collected and processed the samples. C.P. performed the bioinformatics analysis of the Illumina data.
594 B.M., G.C., M.J., J.V.D., L.D., and C.P. drafted the manuscript. All authors read and approved the final
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596 We declare that we have no competing interests.

597

598

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