1 Cooperation and cheating orchestrate *Vibrio* assemblages and

2 polymicrobial synergy in oysters infected with OsHV-1 virus

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- Daniel Oyanedel^{1*}, Arnaud Lagorce^{1*}, Maxime Bruto^{2,3}, Philippe Haffner¹, Amandine Morot^{4,5},
 Yann Dorant¹, Sébastien de La Forest Divonne¹, François Delavat⁶, Nicolas Inguimbert⁷, Caroline
 Montagnani¹, Benjamin Morga⁸, Eve Toulza¹, Cristian Chaparro¹, Jean-Michel Escoubas¹, Yannick
 Labreuche², Yannick Gueguen^{1,9}, Jeremie Vidal-Dupiol¹, Julien de Lorgeril^{1,10}, Bruno Petton^{2,4},
 Lionel Degremont⁸, Delphine Tourbiez⁸, Léa-Lou Pimparé¹, Marc Leroy¹, Océane Romatif¹, Juliette
 Pouzadoux¹, Guillaume Mitta^{1,11}, Frédérique Le Roux^{2,3}, Guillaume M. Charrière¹⁺, Marie-Agnès
- 10 Travers¹⁺, Delphine Destoumieux-Garzón¹
- 11
- 12 ¹ IHPE, Université de Montpellier, CNRS, Ifremer, Université de Perpignan. Montpellier, France
- ² Ifremer, Unité Physiologie Fonctionnelle des Organismes Marins, ZI de la Pointe du Diable,
 Plouzané, France.
- 15 ³ Sorbonne Université, UPMC Paris 06, CNRS, UMR 8227, Integrative Biology of Marine Models,
- 16 Station Biologique de Roscoff, Roscoff cedex, France.
- ⁴ Université de Bretagne Occidentale, CNRS, IRD, Ifremer, LEMAR, F-29280 Plouzané, France
 ⁵ Université de Bretagne-Sud, EA 3884, LBCM, IUEM, F-56100 Lorient, France
- 19 ⁶ Nantes Université, CNRS, US2B, UMR6286, F-44000 Nantes, France
- ⁷ CRIOBE, USR EPHE-UPVD-CNRS 3278, Université de Perpignan Via Domitia, 58 Avenue Paul
 Alduy, 66860 Perpignan, France.
- 22 ⁸ Ifremer, ASIM, Adaptation Santé des invertébrés, La Tremblade, France
- 23 ⁹ MARBEC, Univ Montpellier, CNRS, Ifremer, IRD, Sète, France
- 24 ¹⁰ Ifremer, IRD, Univ Nouvelle-Calédonie, Univ La Réunion, ENTROPIE, F-98800, Nouméa,
- 25 Nouvelle-Calédonie, France.
- 26 ¹¹ Ifremer, Université de Polynésie Française, IRD, ILM, EIO, Polynésie Française.
- 27
- 28 *,+ Equal contribution
- 29 Corresponding authors: Marie-Agnès Travers (Marie.Agnes.Travers@ifremer.fr), Delphine
- 30 Destoumieux-Garzón (delphine.destoumieux-garzon@cnrs.fr)
- 31

32 Abstract

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34 Polymicrobial diseases significantly impact the health of humans and animals but remain understudied in natural systems. We recently described the Pacific Oyster Mortality Syndrome 35 36 (POMS), a polymicrobial disease that impacts oyster production and is prevalent worldwide. 37 Analysis of POMS-infected oysters on the French North Atlantic coast revealed that the disease involves co-infection with the endemic ostreid herpesvirus 1 (OsHV-1) and virulent bacterial 38 39 species such as Vibrio crassostreae. However, it is unknown whether consistent Vibrio populations 40 are associated with POMS in different regions, how Vibrio contribute to POMS, and how they interact with the OsHV-1 virus during pathogenesis. 41

42 We resolved the Vibrio population structure in ovsters from a Mediterranean ecosystem and investigated their functions in POMS development. We find that Vibrio harveyi and Vibrio 43 rotiferianus are the predominant species found in OsHV-1-diseased oysters and show that OsHV-44 1 is necessary to reproduce the partition of the Vibrio community observed in the field. By 45 characterizing the interspecific interactions between OsHV-1, V. harveyi and V. rotiferianus, we 46 47 find that only V. harvevi synergizes with OsHV-1. When co-infected, OsHV-1 and V. harvevi behave 48 cooperatively by promoting mutual growth and accelerating oyster death. V. harveyi showed high 49 virulence potential in oysters and dampened host cellular defenses, making oysters a more 50 favorable niche for microbe colonization. We next investigated the interactions underlying the cooccurrence of diverse Vibrio species in diseased oysters. We found that V. harveyi harbors genes 51 responsible for the biosynthesis and uptake of a key siderophore called vibrioferrin. This 52 important resource promotes the growth of *V. rotiferianus*, a cheater that efficiently colonizes 53 54 oysters during POMS without costly investment in host manipulation nor metabolite sharing.

By connecting field-based approaches, laboratory infection assays and functional genomics, we
have uncovered a web of interdependencies that shape the structure and function of the POMS
pathobiota. We showed that cooperative behaviors contribute to synergy between bacterial and

58	viral co-infecting partners. Additional cheating behaviors further shape the polymicrobial
59	consortium. Controlling such behaviors or countering their effects opens new avenues for
60	mitigating polymicrobial diseases.

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- 62 **Keywords:** Microbiota; mollusk; polymicrobial infection; pathogen; dysbiosis; iron uptake;
- 63 public good; synergy; cooperation; cheating.

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65 Introduction

A number of polymicrobial diseases impact human and animal species [1]. They are defined as 66 67 diseases that result from infections by multiple pathogens [2]. Complex microbe communities that 68 form a cohesive entity with the potential to cause disease in polymicrobial diseases can be referred to as a "pathobiota" [3]. Within pathobiota, microbes synergize to cause disease: their 69 70 interactions enhance disease progression compared to infection with the single microbes [1]. Fatal polymicrobial synergy has been reported between viruses and bacteria including influenza 71 A virus and *Streptococcus* pneumonia [4, 5] or Human Immunodeficiency Virus (HIV) 72 and *Mycobacterium tuberculosis* [6-8]. Synergy has also been reported between viruses such 73 74 as Herpes virus simplex and HIV [9, 10]. Pathobiota show cell-level cooperative capacities that 75 include the production of public goods (e.g. shared metabolites), division of labor, resource 76 transport, and creation and maintenance of the extracellular environment, as in other examples 77 of multicellular organization [11]. Dissecting the interactions within pathobiota is needed to understand and possibly control disease establishment, progression, and symptoms. Theoretical 78 models have been developed in response to these challenges to predict what microbial behaviors 79 80 are favored during such complex interactions [12, 13]. In addition, a series of animal models (both vertebrates and invertebrates) have been used to mimic polymicrobial diseases and validate 81 82 theoretical assumptions [1]. Still, natural pathobiota remain poorly explored.

83 We recently described a typical example of a polymicrobial disease, the Pacific Oyster Mortality Syndrome (POMS). This disease is caused by the Ostreid herpesvirus OsHV-1 and opportunistic 84 bacteria [14] and has devastating consequences for the aquaculture of *Crassostrea gigas* oysters 85 worldwide. The bacterial genera colonizing oysters during POMS are conserved across 86 environments suggesting functional complementarity within the pathobiota [15]. Members of the 87 *Vibrionaceae* family are the best characterized bacteria in the POMS pathobiota [16-20]. Several 88 89 Vibrio species have been shown to have virulence functions in this disease [19, 21]. Vibrio 90 crassostreae (Splendidus clade) uses cytotoxicity and other mechanisms to evade oyster cellular

immune responses, leading to systemic infection [19, 21]. This virulent Vibrio species is highly 91 92 prevalent in OsHV-1-infected oysters on the French Atlantic coast, often associated with other Vibrio species from the superclade Splendidus [17]. A number of studies suggest that Vibrio 93 species found in OsHV-1-infected oysters vary worldwide [22-24]. These species include Vibrio 94 harveyi (superclade Harveyi). A strain of Vibrio harveyi was isolated in 2003 from C. gigas oyster 95 96 spat during a mortality episode in the Thau lagoon (Mediterranean sea) and is pathogenic to oysters [25]. However, we still know little about Vibrio harvevi in POMS and whether V. 97 98 crassostreae or other populations colonize OsHV-1-infected oysters in different ecosystems. In 99 addition, how Vibrio colonizes oysters and how Vibrio interacts with the OsHV-1 virus during 100 pathogenesis remains unclear.

101 Here we performed an integrative study of POMS in a Mediterranean ecosystem, combining field 102 analysis of the Vibrio population structure in OsHV-1-infected oysters with validation of this 103 polymicrobial assembly in mesocosm experiments. We find that two species of the Harveyi clade 104 - namely Vibrio harvevi and Vibrio rotiferianus - are prevalent in diseased oysters in a major 105 Mediterranean area used for oyster farming, the Thau lagoon. Using mesocosm experiments, we 106 characterized the complex interactions between *Vibrio* species and the OsHV-1 virus as well as between the different Vibrio species that assemble in OsHV-1-infected oysters. Our data indicate 107 108 that OsHV-1 infection favors stable colonization by V. harveyi and V. rotiferianus but not other 109 Vibrio of the Harveyi clade. Polymicrobial synergy, including mutual growth promotion and 110 accelerated disease progression, was measured between OsHV-1 and V. harveyi. We next tested 111 the contribution of each partner to the interaction. A series of functional assays, including gene knockouts, indicate that strains of V. harveyi are cytotoxic to immune cells and produce 112 113 siderophores. Our results uncover multiple interdependencies within the POMS pathobiota leading to polymicrobial synergy and accelerated disease progression. We find that initial 114 115 infection with OsHV-1 shapes Vibrio assemblages within the host and favors colonization by V. 116 harveyi. This colonization promotes the growth of OsHV-1 and V. rotiferianus in oysters by

dampening host defenses and by producing vibrioferrin, a key siderophore required for *Vibrio*growth in iron-poor environments.

119 Materials and methods

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121 Oyster and seawater sampling

Juvenile *C. gigas* (pathogen-free diploid oysters produced in hatchery, 6 months old) were immersed in the Thau lagoon (Occitanie, France) and sampled at four timepoints between October 2015 and March 2017. Sampling coincided with mortality events and in the absence of any observed mortality (see Table S1 and S2 for details). 40L of seawater was collected and sizefractionated by sequential filtration (from > 60 µm to 0.2 µm) as described by Bruto et al. [17].

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128 Bacterial sampling

129 Oysters and large seawater particle fractions (> 60 μ m) were ground up with Ultra-Turrax (IKA) and 100 µL was plated on *Vibrio* selective media (thiosulfate-citrate-bile salts sucrose agar, TCBS). 130 Filters of 5, 1 and 0.22 µm porosity were directly placed on TCBS agar and incubated at 20 °C for 131 132 2 days. About 100 colonies per sample were randomly picked then re-streaked first on TCBS and then on Zobell agar (4 g.L-1 bactopeptone, 1 g.L-1 yeast extract and 15 g.L-1 agar in sterile seawater, 133 134 pH 7.4). Stock cultures were stored at -80 °C in Zobell containing 15% glycerol (v/v). For subsequent molecular analyses, all isolates were grown overnight at 20 °C in liquid Zobell medium 135 and bacterial DNA was extracted using the Nucleospin tissue kit following the manufacturer's 136 137 instructions (Macherey-Nagel).

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139 **Population structure analysis**

140 Isolates from October 2015 were genotyped by partial *hsp60* sequencing [26, 27] (Table S3) 141 generating a total of 437 *hsp60* sequences [28]. *hsp60* sequence ambiguities were corrected using 4 peaks and Seaview software (http://nucleobytes.com/index.php/4peaks; [29]) and received a 142 143 taxonomic affiliation if the best BLAST-hit displayed an identity greater than 95% with a type-144 strain. Fisher-exact tests were performed with a 2x2 contingency table using the computing 145 environment R [30] for statistical validation of the ecological preferences of populations and the 146 distribution of bacterial populations in oyster tissues and seawater. Significance was assessed using p-value ≤ 0.05 . 147

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149 MLSA genotyping

150 To validate *hsp60* sequence-based taxonomic assignments to the Harveyi clade, 3 additional 151 protein-coding genes were sequenced (*rctB*, *topA* and *mreB*). First, Harveyi isolates were screened by PCR using rctB-F/rctB-R primers designed to specifically hybridize to Harveyi-related rctB 152 153 gene sequences (Table S3). The PCR program was: 2 min at 95 °C; 30 cycles of 30 sec at 95 °C, 1 154 min at 53 °C and 1.45 min at 72 °C; 5 min at 72 °C. As a result of this analysis, 143 rctB+ isolates 155 were considered to belong to the Harveyi clade. Then *topA* and *mreB* sequences were amplified using VtopA400F/VtopA1200R and VmreB12F/VmreB999R primers, respectively [31, 32], Table 156 S3). All three genes were amplified using the Gotaq G2 flexi polymerase (Promega) following the 157 manufacturer's instructions and sequenced using the reverse PCR primer at GATC Biotech. Next, 158 hsp60, rctB, topA and mreB sequences were aligned with 8 Harveyi superclade type strains using 159 160 Muscle [33]. Alignments were concatenated with Seaview [29]. Phylogenetic trees for each marker were reconstructed with RAxML using a GTR model of evolution and Gamma law of rate 161 162 heterogeneity. Bootstrap values were calculated for 100 replicates. All other options were left at 163 default values.

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165 Bacterial growth conditions

Bacteria were grown for 18 h under shaking at 20°C in Zobell liquid medium (4 g.L⁻¹ bactopeptone, 167 1 g.L⁻¹ yeast extract in sterile seawater, pH 7.4) or LB broth adjusted to 0.5M NaCl unless otherwise 168 stated. When necessary, antibiotics were added (Trimethoprim Trim 10 μ g/mL or 169 Chloramphenicol Cm 10 μ g/mL).

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171 Virulence potential of Vibrio strains

To test virulence potential, Vibrio were grown under shaking at 20 °C for 18 h in Zobell liquid 172 medium before adjustment to $OD_{600} = 0.7$. A volume of 40 µL was injected intramuscularly into 20 173 specific pathogen-free (SPF) juvenile *C. gigas* oysters [16] previously anesthetized in hexahydrate 174 175 MgCl₂ (50 g.L⁻¹, 100 oysters/liter). An injection of *V. crassostreae* J2-9 (virulent strain), *V. tasmaniensis* LMG20012^T (non-virulent strain) or sterile filtered seawater (negative control) were 176 used as controls. After injection, animals were transferred to aquaria (20 oysters per 1 L 177 178 aquarium) containing 400 mL of aerated seawater at 20 °C and kept under static conditions. 179 Mortalities were recorded 24 h post injection.

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181 In vitro cytotoxicity assays

Hemocytes were plated in 96 well-plates (2 x 10⁵ cells/well) as previously published [34]. After 1
h, plasma was removed and 5 μg/μL Sytox Green (Molecular Probes) diluted in 200 μL sterile
seawater was added to each well. Washed *Vibrios* that had been opsonized in plasma for 1 h were
then added to the wells at an MOI of 50:1. Sytox Green fluorescence was monitored (λex 480
nm/λem 550 nm) for 15 h using a TECAN microplate reader. Maximum cytolysis was determined
by adding 0.1% Triton X-100 to hemocytes. Statistical analysis was performed using one-way

- 188 ANOVA and a post-hoc Tukey test for pairwise comparison of maximum cytolysis values for each 189 condition. Significance was assessed using p-value ≤ 0.05 .
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191 Fluorescence microscopy

192 Hemocytes were plated onto glass coverslips in a 24-well plate to obtain monolayers of 5×10^5 cells per well. Adherent hemocytes were exposed to GFP or mCherry-expressing Washed Vibrios 193 194 that had been opsonized in plasma for 1 h, were then added to the wells at a MOI of 50:1. *Vibrios* 195 (Table S4) were added at a multiplicity of infection of 50:1, as in [34]. Binding of bacteria to 196 hemocytes was synchronized by centrifugation for 5 min at 400 g. After a 2 h incubation, the cell 197 monolayers (coverslips from bottom of the wells) were fixed with 4 % paraformaldehyde for 15 198 min. Coverslips were then washed in PBS and stained with 0.25 μ g.mL⁻¹ DAPI (Sigma) and 0.5 µg.ml⁻¹ Phalloidin-TRITC or FITC (Sigma). Fluorescence imaging was performed using a Zeiss 199 200 Axioimager fluorescence microscope and a Zeiss 63× Plan-Apo 1.4 oil objective equipped with a 201 Zeiss MRC black and white camera for image acquisition.

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203 Experimental infection in mesocosm

For experimental infections, a biparental family of oyster (*C. gigas*) spat was produced at the 204 205 Ifremer facilities in La Tremblade (Charente-Maritime, France). This family was selected for its 206 susceptibility to OsHV-1 infection. Spawn occurred in June 2017, and larval and spat cultures were 207 performed as described by Dégremont et al. [35] and Azéma et al. [36]. All growth steps involved 208 filtered and UV-treated seawater. Prior to the experiment, spat were acclimated via a constant 209 flow of filtered and UV-treated seawater enriched in phytoplankton (Skeletonema costatum, Isochrysis galbana, and Tetraselmis suecica) in 120 L tanks at 19°C for at least 2 weeks. Oysters 210 (10 months, 4 cm) were infected with OsHV-1 virus, Vibrio, or both. Microorganisms were 211 212 prepared as follows.

Viral inoculation - For Design 1 (Fig. S1), seawater containing OsHV-1 virions was produced to 213 214 infect pathogen-free juvenile oysters. Briefly, 90 donor oysters were anesthetized, and their adductor muscles were injected with 100 μ l of 0.2 μ m filtrated viral suspension (10⁸ genomic units 215 216 mL⁻¹). These donor oysters were then placed in a 40 L tank for 24 h. Virion release into the 217 seawater was quantified by qPCR. This OsHV-1-contaminated seawater was used to fill tanks for 218 the different experimental conditions. At day 0, 10 recipient oysters were placed in tanks 219 containing 2 L of OsHV-1-contaminated seawater and were sampled at 4, 24, and 48 h; 15 220 additional recipient ovsters were placed in tanks with 3 L of contaminated seawater to track 221 mortalities daily. An identical design was used for control tanks, in which clean seawater was used 222 instead of OsHV-1-contaminated water. For Design 2 (Fig. S1), 250 donor oysters were injected with a filtrated viral suspension as described above. After 24 h, 250 recipient oysters were placed 223 224 in contact with donor oysters in a 40 L-tank. After another 18 h (day 0), recipient oysters were 225 transferred into clean seawater for mortality recording (10 animals in 0.5 L) or for sampling (30 226 animals in 1.5 L).

Bacterial inoculation – Seawater tanks were inoculated with bacteria on day 0 (final concentration of 10^7 CFU/mL). Briefly, bacterial cultures (Zobell broth, 20° C, 18h) were centrifuged at 1500 x g for 10 min. Bacterial pellets were rinsed and resuspended in sterile seawater and the concentration was adjusted to $OD_{600} = 1$ ($10^{\circ} \text{ CFU/mL}$). The bacterial concentration was confirmed by conventional dilution plating and CFU counting on Zobell agar.

At each sampling time point, oysters were sampled together with 100 mL of seawater. The oyster flesh was removed from the shell, snap-frozen in liquid nitrogen and stored at -80°C. 30 mL of seawater were filtered (0.2 μm pore size) and filters were stored at -80 °C. For tissue grinding, individual frozen oysters were shaken for 30 s inside a stainless steel cylinder containing a stainless-steel ball cooled in liquid nitrogen in a Retsch MM400 mixer mill. The pulverized tissue was transferred to a 2 mL screw-capped tube and stored at -80°C until further processing.

238

239 Nucleic acid extraction

Total DNA was extracted from either 20 mg of frozen oyster tissue-powder, 25 mg frozen oyster 240 241 tissue, a pellet from 1 mL of stationary phase bacterial cultures, or a 0.2 µm filter using the 242 Nucleospin tissue DNA extraction kit (Macherey Nagel, ref: 740952.250) with a modified protocol. Briefly, samples were added to a 2 mL screw-capped tube containing Zirconium beads, lysis 243 244 buffer, and proteinase K and shaken for 12 min at a frequency of 35 cycles/s in a Retsch MM400 mixer mill at room temperature and then incubated for 1h 30 min at 56 °C. The samples were then 245 treated with RNase for 5 min at 20 °C and then 10 min at 70 °C. The following purification steps 246 247 were carried out according to manufacturer's recommendations. Total RNA was extracted from 20 mg of frozen oyster tissue-powder using Direct-zol RNA extraction kit (Zymo research). In an 248 249 extra step, the aqueous phase was recovered from the TRIzol reagent prior to column purification 250 as described by [37]; the following steps were carried out as recommended by the manufacturer. 251 Nucleic acid concentration and purity was assayed using a Nanodrop ND-10000 spectrophotometer (Thermo Scientific) and RNA integrity analyzed by capillary electrophoresis 252 on the BioAnalyzer 2100 system (Agilent). 253

254

255 Vibrio genome sequencing and assembly

256 Individual genomic libraries were prepared from 1 ng of bacterial DNA at the Bio-Environment platform (University of Perpignan) using the Nextera XT DNA Library Prep Kit (Illumina) 257 258 according to the manufacturer's instructions. The quality of the libraries was checked using High 259 Sensitivity DNA chip (Agilent) on a Bioanalyzer. Pooled libraries were sequenced in 2x150 paired-260 end mode on a NextSeq 550 instrument (Illumina). Reads were assembled *de novo* using Spades 261 software. Computational prediction of coding sequences together with functional assignments and comparative genomics were performed using the MaGe MicroScope [38]. The genome 262 sequence assemblies have been deposited in the European Nucleotide Archive (ENA) at EMBL-263 EBI under project accession no. PRJEB49488 (Table S6). 264

265

266 **OsHV-1 detection and quantification**

The ground ovster flesh (1.5 mL) was centrifuged for 10 min at 4°C and 2,200 g and genomic DNA 267 was extracted from 50 µL of supernatant using phenol:chloroform:isoamyl alcohol (25:24:1) and 268 isopropanol precipitation. Detection and quantification of OsHV-1 µVar DNA was performed using 269 quantitative PCR targeting a predicted DNA polymerase catalytic subunit (DP) using 270 271 OsHVDPFor/OsHVDPRev primers (Table S3) [39] using the protocol previously described by [37]. 272 All amplification reactions were performed in duplicate using a Roche LightCycler 480 Real-Time 273 thermocycler (qPHD-Montpellier GenomiX platform, Montpellier University). Oyster flesh 274 samples exhibiting a viral load greater than 100 genome units per ng of total DNA (GU/ng) were considered to be infected by OsHV-1. 275

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277 Vibrio quantification

16S rDNA sequences were used to quantify total *Vibrio* present in oysters by extracting 25 ng of
DNA from tissue or crude extracts from seawater (see above). Amplification reactions were
carried out in duplicate, in a total volume of 20 μl on Mx3005 Thermocyclers (Agilent) using
Brilliant III Ultra-Fast SyberGreen Master Mix (Agilent), and 567F and 680R primers at 0.3 μM,
[40], Table S3. Absolute quantification of *Vibrio* genomes in oyster samples was estimated using
standards from 10² to 10⁹ genome copies of *Vibrio* (see supplementary material and methods).

V. harveyi-V. rotiferianus and *V. owensii-V. jasicida* were quantified based on 25 ng of DNA
extracted from tissue or crude extracts from seawater (see above) through detection of a specific
chemotaxis protein and *ompA*, respectively (Table S3). As described above, amplification
reactions were performed in duplicate using a Roche LightCycler 480 Real-Time thermocycler,
SYBR Green I Master mix (Roche) and primers at 0.3 and 0.2 μM f.c. respectively. For absolute

quantification, standard curves of known concentration of *V. harveyi*, *V. rotiferianus*, *V. jasicida*, *V. owensii* genomes were used (supplementary material and methods).

291

292 *rctB* metabarcoding

Locus-specific PCR primers, including Illumina overhang adaptors, were designed to amplify a 293 294 573 bp region of the *rctB* gene in all our *Harvevi* strains, (rctB-Fw-I and rctB-Ry-I primers, Table 295 S3). PCR analysis of total DNA extracted from oysters (N=60) used the high fidelity Q5 polymerase 296 (New England Biolabs) in a total volume of 50 μ L under the following conditions: 98 °C for 25 s 297 followed by 35 cycles of 98 °C for 10 sec, 51 °C for 25 sec and 72 °C for 30 sec. Final extension was performed at 72 °C for 2 minutes. Presence of the 573 bp amplicon was validated by 1.5 % gel 298 299 electrophoresis. Libraries were constructed with the Two-Step amplicon sequencing approach 300 using Illumina dual indexes (ref. 15044223) and sequenced on a MiSeq instrument to produce paired end reads 2x300 bp, by the GenSeq platform, University of Montpellier (ISEM), France. 301 302 Sequencing data were processed using the SAMBA pipeline v3.0.1. [41]. All bioinformatics processes used the next-generation microbiome bioinformatics platform QIIME 2 [42] (version 303 2020.2) and grouped sequences in ASV (Amplicon Sequence Variants) using DADA2 v1.14 [43]. 304 305 The resulting ASVs were annotated against an in-house database containing the Harveyi *rctB* 306 sequences and filtered for low abundance ASVs to limit the prevalence of putative artifacts due to 307 sequencing errors. To do this, we only retained ASVs showing at least four reads in at least four 308 samples. Statistical analyses were performed with R [30] using the R packages Phyloseq v1.38.0 [44] and Vegan v2.6-2 [45]. Principal coordinate analyses (PCoA) based on Bray-Curtis distances 309 310 at each kinetic point were used to assess variation in the composition of Harveyi communities. Putative differences between groups were assessed by statistical analyses (Permutational 311 312 Multivariate Analysis of Variance - PERMANOVA) using the function adonis2 implemented in 313 vegan [45]. Finally, we used DESeq2 v1.36.0 and STAMP software [46] to identify ASVs with significant variation in abundance. 314

315

316 Mutagenesis

Deletion of pvuA1 (THOG05 v1 100041) and pvuA2 (THOG05 v1 100042) in V. rotiferianus 317 Th15_0_G05 was achieved through double homologous recombination between the pLP12 318 suicide plasmid (Table S8) and the bacterial chromosome [47]. Briefly, two fragments of around 319 320 800 bp flanking the target region were amplified, assembled by GeneArt, and cloned into the 321 pLP12 plasmid [48]. The suicide plasmid (named pAM010) was transferred by conjugation 322 between an *Escherichia coli* β3914 donor [49] and *V. rotiferianus* Th15_0_G05 recipient using a 323 triparental mating procedure (Table S4-5). The first and second recombination events leading to 324 pAM010 integration and elimination were selected following a recently published method [47]. Mutants were screened by PCR using primers del-pvuA1-A2-OG05-F and del-pvuA1-A2-OG05-R 325 (Table S3). A V. rotiferianus Th15_0_G05 mutant strain deleted for the pvuA1-2 genes was stored 326 in glycerol at -80 °C (strain *V. rotiferianus* Th15_0_G05 Δ*pvuA1-2*). 327

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329 Vibrio growth in iron-depleted media and rescue

Growth experiments in iron-depleted medium were performed with *V. rotiferianus* Th15_0_G05 330 331 and its $\Delta pvuA1-2$ isogenic derivative. Isolates were grown overnight at room temperature in 332 Artificial Sterile Seawater (ASW: NaCl 40 mM, KCl 20 mM, MgSO₄ 5 mM, CaCl₂ 2 mM) with 0.3 % 333 (wt/vol) casamino acids and vitamins (0.1 μ g/L vitamin B12, 2 μ g/L biotin, 5 μ g/L calcium 334 pantothenate, 2 µg/L folic acid, 5 µg/L nicotinamide, 10 µg/L pyridoxin hydrochloride, 5 µg/L riboflavin, 5 mg/L thiamin hydrochloride). Cultures were pelleted (2 min at 15,000 g) and washed 335 in ASW. Cells were then inoculated (1:100) into minimal media with (iron-poor) or without (iron-336 replete) the iron-specific chelator 2,2'-bipyridin (150 µM). Bacteria were grown in 96-well plates 337 with orbital shaking in a Tecan microplate reader (Infinite M200) at 25 °C for 24 h. The OD₆₀₀ was 338 339 recorded at 30 min intervals. Rescue was performed either by adding freeze-drying concentrated

V. harveyi Th15_0_G11 cell-free supernatant or by adding synthetic vibrioferrin (8.7 to 70 μM) to
cultures. Vibrioferrin was synthesized according to Takeuchi et al. [50](Fig. S9). NMR and HRMS
spectroscopic data confirmed the structure (Fig. S10) and were consistent with the literature [50].

343

344 **Results**

345 Specific Vibrio populations assemble in oysters infected by OsHV-1 virus

We asked whether Vibrio populations that naturally assemble in oysters infected by OsHV-1 were 346 conserved across farming environments. With this objective in mind, we performed a field 347 348 experiment in the Thau lagoon (South of France). *Vibrio* populations have been suggested to be 349 distinct in this Mediterranean ecosystem from those previously found in the Atlantic (bay of Brest, 350 northwest of France) [25]. We performed our study during an episode of oyster mortality. Specific 351 pathogen-free (SPF) juvenile oysters were immersed in September 2015 in the Thau lagoon, 352 which hosts significant oyster farming activity (352 acres). Oysters tested positive for OsHV-1 353 after one month (October 2015), indicating an ongoing episode of POMS (Table S1). We 354 characterized the population structure of the Vibrio isolated from a pool of oysters and from the 355 surrounding seawater. A total of 472 isolates were sampled on Vibrio-selective medium from infected oyster tissues and from the water column (Table S2). Partial hsp60 sequences were 356 obtained for 437 isolates. Assignment to the Vibrio genus was confirmed for 304 sequences (67.8 357 %), which exhibited \geq 95 % identity with the *hsp60* sequence from a *Vibrio* type-strain (Table S2)., 358 359 Isolates with *hsp60* sequence identity below this threshold were not included in the study. We 360 observed contrasting population structures in oyster tissues and in the water column. Oyster 361 tissues were dominated by 3 populations of the Vibrio Harveyi super clade (V. harveyi, V. 362 rotiferianus and V. owensii), representing 54/55 isolates (Fig. 1A). The water column showed a higher diversity of Vibrio with only 53/249 isolates (21%) falling into the Harveyi super clade 363 (Fig. 1A, Fig. S2). These water column isolates were dominated by *V. jasicida* (39/53) (Fig. 1A). 364 Taxonomic assignments of Harveyi-related isolates were confirmed by multi-locus sequence 365

analysis (MLSA) phylogeny based on 4 Vibrio genes (hsp60, rctB, topA and mreB) (Fig. 1B) 366 367 (Lagorce, 2022). A total of 101 isolates could be assigned to V. harveyi (n=63), V. rotiferianus (n=17), V. jasicida (n=15) and V. owensii (n=6) (Fig. 1B). Among them, V. harveyi and V. rotiferianus 368 369 showed a positive association with oyster tissues (Fisher exact test, p < 0.001) whereas V. jasicida 370 was almost exclusively associated with the water column (Fisher exact test, p < 0.001) (Fig. 1B). 371 The high relative abundance of *V. harveyi* in POMS-diseased oysters from the Thau lagoon was 372 confirmed by three subsequent samplings in 2016-2017: V. harveyi was isolated during but not outside of POMS episodes, almost exclusively from ovsters infected with OsHV-1 (Table S1). This 373 374 result contrasts with the preferential association of the species V. crassostreae with POMS-375 diseased oysters on the French Atlantic coast [17].

376

377 Preferential association of V. harveyi and V. rotiferianus with OsHV-1-infected oysters

378 As a number of interdependent environmental variables (e.g. temperature, salinity, season, viral 379 infection) may influence Vibrio assemblages in oysters in the field, we next tested how Vibrio 380 populations partition and the role of OsHV-1 in this partitioning. To achieve this aim, we used 381 mesocosms, which allow controlled infection experiments of oysters through natural routes with 382 microorganisms relevant to POMS. We developed a synthetic Vibrio community composed of a 383 mixture of 20 isolates from the Harveyi super clade in the presence (VO) or absence (V) of OsHV-384 1 (Fig. 2A). Vibrio representative of the 4 populations isolated from the Thau lagoon were used: V. harveyi and V. rotiferianus (positively associated with oysters), V. jasicida (negatively associated 385 386 with oysters) and *V. owensii* (neutral). In parallel, oysters were exposed to OsHV-1 virus only (O) or were kept in tanks devoid of introduced pathogens (control). Oysters were collected in the first 387 388 48 h, before mortalities occurred. We first examined which Vibrio populations colonized oysters 389 in the presence/absence of the OsHV-1 virus by comparing the V and VO conditions. To 390 discriminate between the four populations introduced in the mesocosm, we developed an 391 amplicon sequencing method based on the *rctB* polymorphic gene (Fig. S3). In the absence of 392 OsHV-1, the Harveyi-related population assemblage remained stable in oysters over time, as 393 shown by *rctB*-barcoding (Fig. S5). In contrast, co-infection with OsHV-1 had a significant effect 394 on the structure of the assemblage, as observed after 48 h (Permutational multivariate analysis of 395 variance, p = 0.001 (Fig. 2B, Fig. S4-S5). The species V. harveyi and V. rotiferianus were 396 significantly enriched in oyster flesh in the presence of 0sHV-1 (p < 0.05, Welch's t-test, p-value 397 corrected with Benjamini-Hochberg FDR) (Fig. 2C). In contrast, V. owensi was equally abundant in 398 the presence/absence of OsHV-1, and *V. jascida* was more abundant in oyster flesh in the absence of OsHV-1 (p < 0.001) (Fig. 2C). The positive effect of OsHV-1 on ovster colonization by V. 399 400 harveyi/V. rotiferianus but not V. jasicida/V. owensii at 48 h was confirmed by qPCR monitoring of pathogen loads (mutiple t test, p < 0.01) (Fig. S4). Altogether, our experimental results show (i) 401 that OsHV-1 is necessary to reproduce the distribution of Vibrio community observed in the field 402 during a POMS episode, and (ii) that only V. harveyi and V. rotiferianus efficiently colonize OsHV1-403 404 infected oysters.

405

406 OsHV-1 virus synergizes with V. harveyi and/or V. rotiferianus in oyster mortality

407 We next tested the effect of the Vibrio/OsHV-1 interaction on POMS progression. We used the same synthetic community of *Vibrio* (Fig. 2A) to monitor oyster mortality and pathogen loads 408 under different experimental conditions. Mortalities were only observed in tanks containing the 409 OsHV-1 virus, indicating that the synthetic Vibrio community alone (V) was not lethal to oysters 410 through natural infection routes. Mortalities started at day 2 in tanks containing OsHV-1 (both O 411 412 and VO conditions). Still, mortalities progressed significantly more rapidly in oyster tanks containing both OsHV-1 and the synthetic Vibrio community (VO) with 90% mortalities in 3 days 413 414 as opposed to 6 days for oysters exposed to OsHV-1 only (O) (Kaplan-Meier survival curves, log-415 rank test, p = 0.0018) (Fig. 2D, left panel). Therefore, introducing the synthetic Vibrio community accelerated the OsHV-1-induced disease. As expected from our previous study [14], the total 416 417 Vibrio load increased constantly over 48 h in OsHV-1-infected oysters (O). Remarkably, the

increase in Vibrio load was significantly higher in oysters exposed to both the Vibrio community 418 419 and OsHV-1 (VO) than in oysters exposed to Vibrio only (O) at 48h, *i.e.* at the onset of mortality 420 (Kruskal-Wallis test, p < 0.001) (Fig. 2D, right panel), with a significant contribution of V. harveyi 421 and V. rotiferianus (Fig. S4). In oysters exposed to Vibrio only (V), i.e. oysters that did not die, Vibrio 422 colonization tended to be transient with a peak between 4 h-24 h (Fig. 2D, right panel). Altogether, 423 this indicates that in the absence of OsHV-1, oysters tolerate transiently high loads of the synthetic 424 Vibrio community but Vibrio colonization is not stable and ultimately decreases without causing mortality. In contrast, OsHV-1 infection favors the proliferation and persistent colonization of 425 426 Vibrio such as V. harveyi and V. rotiferianus, which exacerbate pathogenesis, an effect not seen 427 with V. jascida and V. owensii. These data show that OsHV-1 and specific populations of Vibrio act 428 in synergy to accelerate oyster death.

429

430 *V. harveyi* and OsHV-1 reciprocally promote inside-host growth

431 To get insight into the synergistic process, we next tested whether OsHV-1 and strains of *V. harveyi* 432 or *V. rotiferianus* affect one another's growth in co-infections. To facilitate pathogen monitoring 433 oysters were exposed to OsHV-1 and fluorescent *Vibrio* strains representing each population (Fig. 434 3A). Infection with OsHV-1 only was used as a control (Fig. 3A). As in the previous mesocosm 435 experiment (Fig 2), live oysters were sampled at 0, 4, 24 and 48 h, before the onset of mortalities 436 (Fig. S6) to monitor pathogen loads in every individual. First, we compared V. harveyi and V. rotiferianus colonization in oysters infected with OsHV-1. Only V. harveyi had the ability to 437 438 colonize OsHV-1-infected oysters efficiently. Indeed, V. harveyi remained present at high doses $(10^5 \text{ to } 5 \times 10^6 \text{ copies/ng of DNA})$ in live oyster tissues throughout the time course. In contrast, V. 439 440 *rotiferianus* loads decreased rapidly over the same period and were undetectable (< 10⁴ copies/ng 441 of DNA) in most individuals after 48 h (Fig. 3B). Second, we analyzed the effect of Vibrio strains on OsHV-1 growth. Remarkably, the viral load was 100-fold higher at 48h in oysters co-infected 442 443 with *V. harveyi* and OsHV-1 than in oysters infected with OsHV-1 only (t-test, p < 0.05). Such an

increase in viral load was not observed in co-infections with *V. rotiferianus* (Fig. 3C), consistent
with the rapid elimination of *V. rotiferianus* from host tissues (Fig. 3D). Altogether, these results
indicate that *V. harveyi* and OsHV-1 cooperate by increasing mutual growth during pathogenesis,
an effect not observed with *V. rotiferianus*.

448

449 V. harveyi actively dampens oyster immune defenses

450 We next investigated Vibrio traits that may facilitate host colonization and favor polymicrobial 451 synergy with OsHV-1. We first compared the virulence potential of V. harveyi and V. rotiferianus 452 (successful colonizers, Fig. 1B, 2C) and V. jasicida and V. owensii (poor colonizers). Virulence 453 potential was tested in vivo by direct injection of Vibrio isolates into oyster adductor muscle. V. 454 harveyi isolates showed strong virulence potential as revealed by an average of 50% oyster 455 mortality one day after injection (Fig. 4A). However, mortalities remained below 15% on average 456 after injection of V. rotiferianus, V. jasicida, and V. owensii (Fig. 4A). Not only did V. harveyi have a 457 significantly higher virulence potential than other species (Kruskal-Wallis test, p < 0.001), but it 458 also showed greater cytotoxicity toward oyster immune cells (hemocytes) in vitro (Fig. 4B). Here 459 we compared the cytotoxic activity of two strains per Vibrio species among the four Harveyi-460 related species from our study. Upon exposure to V. harvevi strains, 67-76% of hemocytes underwent lysis (p < 0.001, One-way ANOVA, and Tukey post-hoc test) (Fig. 4B). The three other 461 462 *Vibrio* species were not cytotoxic, with the percentage of lysed cells similar to controls (7 to 23%). We further confirmed the ability of *V. harveyi*, but not *V. rotiferianus*, to damage hemocytes using 463 464 fluorescent Vibrio strains. After a 2 h exposure to V. harveyi, hemocytes were massively damaged 465 and many extracellular V. harveyi were observed (Fig. 4C). In contrast, no cellular damage was observed when hemocytes were exposed to V. rotiferianus, with most bacteria being phagocytized. 466 467 Thus, V. harveyi strains are equipped with specific virulence/cytotoxicity mechanisms that can dampen oyster cellular defenses. To uncover genomic features that might explain V. harveyi 468 469 virulence/cytotoxicity, we sequenced and analyzed 17 Harveyi-related genomes from our

collection (4 *V. harveyi*, 5 *V. owensii*, 4 *V. jasicida* and 4 *V. rotiferianus* isolates; Table S6). We found
that the *V. harveyi* genome contained the most candidate virulence factors of the four *Vibrio*species in the present study. Candidate virulence genes in the *V. harveyi* genome included a type
3 secretion system (T3SS) and its associated effectors as well as 3-4 different type 6 secretion
systems (T6SS) (Table S7). Such T6SS were previously shown to be essential for oyster
colonization in *V. crassostreae* and *V. tasmaniensis* [19, 21].

476

477 *V. harveyi* produces vibrioferrin, which can be used by *V. rotiferianus*

478 We next used comparative genomics to examine how V. harveyi and V. rotiferianus both colonize 479 OsHV-1-infected ovsters but make distinct contributions to pathogenesis (Table S6). Using genome comparisons, 15 genes were identified that are exclusively shared by V. harvevi and V. 480 481 rotiferianus (good colonizers; Table S8). Of these, iron acquisition systems differed between species (Fig. S7). We paid particular attention to genes involved in the vibrioferrin pathway, a 482 siderophore whose uptake system was only found in strains of V. harveyi and V. rotiferianus (Fig. 483 484 5A). Vibrioferrin is a tricarboxylic acid siderophore derived from citric acid; it was shown to be shared as a public good within populations of *V. splendidus* [51]. Remarkably, we found that *V.* 485 harveyi harbors genes responsible for the biosynthesis and uptake of vibrioferrin whereas V. 486 487 rotiferianus only carries the vibrioferrin uptake system (Fig. 5A). This suggests that by secreting 488 vibrioferrin, V. harveyi could facilitate iron uptake by V. rotiferianus. To determine whether V. 489 rotiferianus is able to use vibrioferrin produced by V. harveyi, we compared the growth of V. 490 rotiferianus in the presence/absence of increasing amounts of V. harvevi culture supernatant. V. 491 *harveyi* was able to grow in iron-depleted medium (2,2'-bipyridine used as iron chelator, Fig. S8), 492 in agreement with its ability to produce siderophores. In contrast, growth of V. rotiferianus was 493 impaired upon iron depletion, (p < 0.05, Mann-Whitney) (Fig. S8). Growth of V. rotiferianus was rescued in a dose-dependent manner by the addition of cell-free culture supernatant from V. 494 495 *harveyi* (Kruskal-Wallis, p < 0.05) (Fig. 5B). To determine whether this rescue is linked to vibrioferrin, we synthesized the siderophore (Fig. S9-S10). Synthetic vibrioferrin was sufficient to 496

497rescue V. rotiferianus growth in iron-depleted medium (Kruskal-Wallis, p < 0.05) (Fig. 5C). Finally,498we constructed a V. rotiferianus mutant strain which lacks the two genes encoding the PvuA1-A2499receptor that are required for vibrioferrin uptake. Vibrioferrin failed to rescue the growth of the500V. rotiferianus $\Delta pvuA1-A2$ mutant (Fig. 5D). This demonstrates that V. rotiferianus is able to501acquire vibrioferrin, an important resource produced by V. harveyi (and potentially other Vibrio502within the microbiota), to grow in iron-poor environments.

Overall, our data show that *V. rotiferianus* behaves as a cheater by using a siderophore produced
by *V. harveyi*. Thus unlike *V. harveyi* and the OsHV-1 virus, which behave cooperatively, *V. rotiferianus* successfully colonizes oysters taking advantage of public goods contributed by the
microbiota without providing benefit to the POMS-associated microbiota.

507

508

509 Discussion

Here, we used a natural pathosystem, the Pacific Oyster Mortality Syndrome (POMS), to disentangle the complex web of interactions that shape polymicrobial assemblages and pathogenicity. Our data highlighted a web of interdependencies in which diverse microorganisms shape the POMS pathobiota and accelerate disease progression. In particular, we identified conserved cooperative traits accelerating disease progression as well as incidental cheating traits. These traits contribute to shaping the *Vibrio* community associated with OsHV-1-infected oysters.

We showed here that the population structure of *Vibrio*, a genus consistently found in the POMS pathobiota [52], varies across French Atlantic and Mediterranean oyster-farming environments. Indeed, we found *V. harveyi* and *V. rotiferianus* (Harveyi clade) positively associate with OsHV-1infected oysters during field mortalities in the Mediterranean. This finding contrasts with a previous analysis of Atlantic oysters where *V. crassostreae* (Splendidus clade) dominated [17]. Experimentally, two out of four different species from the same clade were recruited from seawater, suggesting that the environment serves as a reservoir for *Vibrio* recruitment during 523 POMS. Differences in Vibrio species associated with OsHV-1-infected oysters are therefore likely 524 related to different environmental distributions of Vibrio species. Indeed, while V. crassostreae is 525 present at high latitudes, e.g. in the North sea, Germany [53], V. harveyi preferentially grows in warm waters [54], such as those found for half the year in the Thau lagoon (16-30°C) [55]. 526 527 Consistently, V. harveyi has been associated with disease outbreaks under rising sea surface 528 temperatures [56]. Remarkably, the data from both locations (Atlantic/Mediterranean) converges 529 in that the OsHV-1 virus is associated with specific populations of *Vibrio* in diseased oysters only. 530 Our data indicate that OsHV-1 is key to enabling colonization by such specific *Vibrio* species.

By focusing on Vibrio that naturally co-infect oysters with OsHV-1, we observed a first level of 531 532 interdependence and polymicrobial synergy occurring during POMS. Indeed, the combined effects 533 of OsHV-1 and Vibrio triggered a faster host death than that observed when the microorganisms 534 were used in isolation to infect oysters. While both V. harveyi and V. rotiferianus successfully colonized OsHV-1-infected oysters, polymicrobial synergy was specifically observed between 535 OsHV-1 and V. harveyi. The benefits of co-infection were not only observed for OsHV-1 but also 536 for the whole Vibrio community: both showed significantly greater expansion prior to oyster 537 538 death when OsHV-1 and V. harveyi were co-infected. Whether other bacterial genera conserved 539 among POMS consortia (e.g. Arcobacter, Marinomonas) [15, 52, 57] also contribute to this 540 polymicrobial synergy or play complementary roles in the polymicrobial consortium remains to be established. 541

A key mechanism underlying polymicrobial synergy between OsHV-1 and *Vibrio* is the dampening 542 of oyster cellular defenses, which makes the local environment less hostile for the entire 543 microbiota. This manipulation of cellular immunity is key in a number of polymicrobial infections. For 544 545 instance, induction of IL-10-producing macrophages by *M. tuberculosis* favors HIV-1 replication and 546 spread [8]. Similarly, the apoptosis of macrophages, neutrophils, dendritic cells and NK cells 547 contributes to the fatal outcome of influenza A virus / S. pneumoniae co-infections [5]. Importantly, 548 cytotoxicity to host immune cells is a phenotypic trait conserved in Vibrio species that co-occurs with 549 OsHV-1 in diseased oysters. Indeed, cytotoxic effects toward oyster hemocytes is a functional trait

conserved in Mediterranean strains of V. harveyi, as well as in strains of V. crassostreae, V. 550 551 tasmaniensis and V. splendidus [19, 21, 58] isolated from POMS-diseased oysters in the Atlantic 552 [17, 20, 25, 59, 60]. We previously showed that cytotoxicity toward immune cells is essential for Vibrio to colonize oyster tissues [21]. This indicates that Vibrio species that harbor similar 553 554 functions are likely replaceable within the POMS bacterial consortia. Our mesocosm experiment 555 validated the preferential association of V. harveyi with OsHV-1-infected oysters under controlled 556 conditions, showing that cytotoxic *Vibrio* species can be recruited from the ecosystems where they 557 circulate. In the mutualistic association between the mollusk *Euprymna scolopes* and its symbiont 558 Vibrio fischeri, it was also shown that Vibrio phenotypic traits determine their capacity to be 559 recruited from the environment [61]. Overall, our data show that similar to OsHV-1 [14], cytotoxic 560 *Vibrio* species modify their extracellular environment by targeting oyster immunity. This results 561 in eased proliferation of co-infecting partners, as shown here for OsHV-1 and specific Vibrio 562 species. This fundamental cooperation taking place within the POMS pathobiota has benefits for 563 the entire polymicrobial consortium.

564 Our present study suggests that cooperation through dampening host defenses contributes to the 565 shaping of the POMS pathobiota assembly. This supports recent studies highlighting that cooperative interactions within animal microbiota participate in the shaping of community 566 567 composition and functioning (for review see [62]). However, pathogenic microbes cooperating to 568 manipulate their host may be outcompeted by other members of the microbiota that do not invest 569 in cooperation. Manipulation is therefore only expected to be favored if its benefits predominantly 570 fall back on the manipulator [63]. One important benefit for OsHV-1 and V. harvevi observed here is a higher load for both manipulators. Another likely benefit is increased pathogen transmission 571 572 due to accelerated host death (Fig. 6). From a general point of view, accelerating host death is not predicted to be favorable to pathogens as this may drive the host, and consequently themselves, 573 574 to extinction, unless transmissibility is also increased [64]. This is particularly true for pathogens 575 with narrow host spectra like V. harveyi and V. crassostreae, but also for OsHV-1, which are almost exclusively found in oysters (this study;[17]). POMS is typically transmitted from oyster to oyster 576

577 by a massive release of pathogens into the seawater, which in turn infects neighboring oysters 578 through filter-feeding [57, 65]. We can therefore hypothesize that polymicrobial synergy leading 579 to accelerated release of OsHV-1 and *Vibrio* into seawater is advantageous in terms of group 580 selection.

Beyond cooperative traits conserved among successful *Vibrio* colonizers, we find evidence that cheating is an efficient strategy that *Vibrio* use to colonize oysters affected by POMS. Indeed, unlike *V. harveyi, V. rotiferianus* does not invest resources in virulence nor cytotoxicity. Moreover, our data argues that *V. rotiferianus* lacks the costly pathways to synthesize vibrioferrin. Instead, this species imports this siderophore, produced by *V. harveyi*, to promote its own growth. In contrast, the unsuccessful colonizers *V. owensii* and *V. jasicida* lack the vibrioferrin uptake machinery, which may contribute to their competitive exclusion [66].

588 Remarkably, vibrioferrin biosynthetic pathways are highly conserved between V. harveyi and V. 589 crassostreae, the two main species associated with POMS in the Mediterranean and the Atlantic. 590 Comparison of these *Vibrio* genomes reveals conserved synteny and > 70% amino acid identity in 591 homologous proteins involved in these pathways (Fig. S12). Both Vibrio species can, therefore, supply similar public goods to the POMS pathobiota. Metabolite cross-feeding, which enables a 592 593 bacterium to consume metabolites produced by another community member, can mediate 594 synergy in multi-species infections (for review see [1]). Cheating behavior regarding iron-595 acquisition was earlier described in Vibrio by Cordero et al. [51]. The authors showed that within 596 ecologically cohesive clusters of closely related *Vibrio*, only some genotypes were able to produce 597 siderophores. Meanwhile, non-producers had selectively lost siderophore biosynthetic pathways. 598 We observe traces of selective loss in *V. rotiferianus* strains: while *V. rotiferianus* and *V. harveyi* both harbor the genomic region that contains vibrioferrin receptors, the former species 599 600 specifically lacks the vibrioferrin biosynthetic genes.

601 The cheating behavior arguably coevolves with ecological adaptation of *Vibrio* toward association602 with larger particles in the water column, consistent with efficient siderophore sharing where

603 local densities of bacteria are high [51]. Because oysters host dense populations of bacteria in 604 their body fluids (> 10^7 culturable bacteria/mL during episodes of POMS [67]), they constitute 605 microhabitats where cheating can occur. Oysters could, therefore, provide a favorable niche for 606 Vibrio strains that can import iron-loaded siderophores. Importantly, the present study shows 607 that cheating for iron acquisition occurs inter-specifically and coincides with the co-occurrence of 608 V. harveyi and V. rotiferianus in POMS-diseased oyster. Indeed, our two-year Mediterranean field 609 survey showed that V. harveyi is repeatedly associated with diseased oysters. V. harveyi co-occurs 610 with *V. rotiferianus* in ovster flesh, and to a lower extent in particles >60µm (zooplankton) and 5-611 60µm (phytoplankton). These host-associated states provide conditions where V. rotiferianus could have evolved cheating behavior. Thus, social interactions through siderophore-sharing 612 appear to structure the assembly of bacterial species in oysters affected by POMS. 613

614

615 **Conclusion**:

616 By disentangling the complex interactions at play in the Pacific Oyster Mortality Syndrome, we 617 have shown that cooperation is key in the functioning of this natural pathosystem. Cooperation 618 and cheating seem to drive the assembly of the pathobiota. The former manifests as polymicrobial 619 synergy between the OsHV-1 virus, the etiological agent of POMS and secondary *Vibrio* colonizers. 620 Dampening of oyster cellular defenses and siderophore sharing, both of which make the host 621 environment more favorable for microbial proliferation, are two cooperative traits conserved in the main POMS-associated Vibrio species, namely V. harveyi and V. crassostreae, across oyster 622 623 farming environments. This knowledge opens new avenues for the control of polymicrobial diseases by interfering with polymicrobial assembly. Implementation of ecological principles, 624 625 such as interfering with cooperative behavior within the microbiome (*e.g.* siderophore sharing) 626 or altering the local environment of the POMS pathobiota (*i.e.* stimulating host defenses) are 627 promising solutions for future exploration. We have shown recently that eliciting oyster antiviral 628 defenses through so-called immune priming is sufficient to prevent colonization by OsHV-1 and

subsequent disease development [37, 68]. Similarly, exposure to microbial communities at early
developmental stages (referred to as biological embedding) was also protective against POMS
[69]. Such solutions, which make the host a less favorable niche for the OsHV-1 virus [68] and/or
bacteria [69], are promising avenues for preventing or reducing the assembly of microbial
consortia that produce POMS.

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- 635

636 **Declarations**

637 *Ethics approval*

The animal (oyster *Crassostrea gigas*) testing followed all regulations concerning animal experimentation. The authors declare that the use of genetic resources fulfill the French regulatory control of access and EU regulations on the Nagoya Protocol on Access and Benefit-Sharing (ABSCH-IRCC-FR-259502-1).

642

643 Availability of data and materials

644 Targeted gene sequences (hsp60, rctB, topA and mreB) and all sequence files with associated metadata generated in mesocosm experiments are available in Ifremer Oceanic database (doi 645 646 doi.org/10.12770/173c0414-a3ca-4a79-b6b2-cd424ee90593 [28] and doi.org/10.12770/63b02659-cd9d-4834-8e6d-8adfa736755d [70]). Genome assemblies have 647 648 been deposited in the European Nucleotide Archive (ENA) under project accession no. PRJEB49488. Original R statistic scripts for metagenomics analyses and the phyloseq table are 649 650 available https://doi.org/10.5281/zenodo.7599486.

651

652 *Competing interests*

653 The authors declare that they have no competing interests.

654

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661

662 Author contributions :

Conception: DDG, MAT, GMC, FLR, GM. Designed the work: DDG, MAT, GMC, FLR, GM, DO, AL.
Acquisition, analysis and/or interpretation of data: DO, AL, MB, PH, AM, YD, FD, NI, BM, ET, CC,
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MAT, GMC, DO, AL or substantively revised it: FLR, GM, JME JDL.

667

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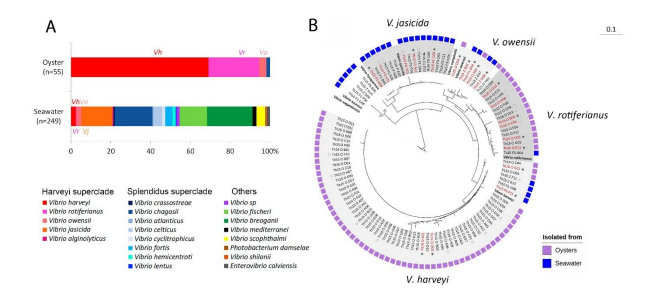


Figure 1. Vibrio harveyi and Vibrio rotiferianus are the most prevalent Vibrio species in OsHV-1-infected oysters

The population structure of *Vibrionaceae* was determined in seawater and oyster flesh during an episode of POMS (Thau lagoon, October 2015). **(A)** shows the distribution (%) of *Vibrio* isolated from oysters and seawater. A total of 304 isolates whose *hsp60* sequence displayed \geq 95% identity with a *Vibrio* type strains were included. A high prevalence of the Harveyi super-clade (*V. harveyi* –*Vh, V. owensii* –*Vo, V. jasicida* –*Vj* and *V. rotiferianus* –*Vr*) is observed in oysters. **(B)** shows the high prevalence of *V. harveyi* and *V. rotiferianus* in oysters after taxonomic affiliation shown in A was validated by an MLST analysis. Four marker genes (*hsp60, rctB, topA* and *mreB*) were used. A phylogenetic tree was constructed with the concatenated sequences of the 4 markers. The following reference strains were used in the analysis: *V. azureus* NBRC 104587, *V. campbellii* CAIM 519, *V. harveyi* NBRC 15634, *V. hyugaensis* 090810a, *V. jasicida* CAIM 1864, *V. owensii* CAIM 1854, *V. rotiferianus* CAIM 577, *V. sagamiensis* NBRC 104589. *Vibrio sagamiensis* type strain was used as an outgroup to root the tree. Bootstrap values (>50%) are indicated by circles on branches. Color boxes indicate whether a strain was isolated from seawater (blue boxes) or from oysters (purple boxes). For details on seawater column fractionation, see Fig. S2. Stars indicate strains whose genome was sequenced in this study, and filled stars those used for comparative genomics. Red indicates strains used in mesocosm experiments (Fig. 2).

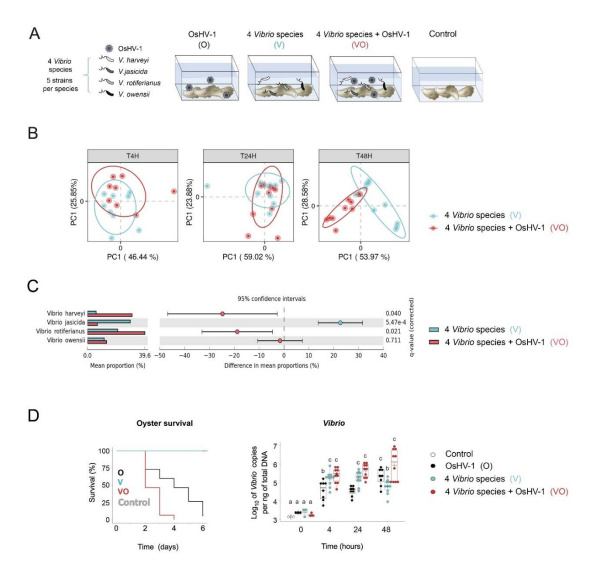


Figure 2. OsHV-1 synergizes with Vibrio harveyi and/or Vibrio rotiferianus to kill oysters

(A) Mesocosm experiment to test OsHV-1 synergy with Vibrio species (Design 1). Specific pathogen-free oysters were placed in contact with seawater containing OsHV-1 (108 genomic units mL-1) or Vibrio (107 CFU.mL-1) or both during 6 days at 20°C. (0): seawater containing only OsHV-1. (V): 4 species of the Harveyi clade (V; V. harveyi, V. rotiferianus, V. owensii, V. jasicida), (VO): both OsHV-1 and Vibrio, (C): Controls not exposed to pathogen. (B) Monitoring of the OsHV-1-induced dysbiosis by rctB-barcoding. Principal Component Analysis (PCoA) ordination plots of Bray-Curtis dissimilarities for the Harveyi-related community associated with oysters. PCoA results are depicted for each time point (i.e. 4, 24 and 48 h). Each dot represents the Vibrio microbiome of one oyster. Colors refer to the experimental condition (blue for V, red for VO). Ellipses represent the 95% confidence intervals for each group. PERMANOVA between the two experimental conditions and the three time-points indicates significant differences p = 0.001 (see Fig S5). Statistical differences between V and VO conditions are observed at 48 h. (C) Colonization of diseased oysters by V. harveyi and V. rotiferianus at 48 h. The left barplot represents the mean proportion of the species V. harveyi, V. rotiferianus, V. owensii, V. jasicida in oysters from the V and VO conditions at T=48h. The right dot plot represents the difference in mean proportion of each species by STAMPS analysis. Statistical differences were obtained from Welch's t-test (p-value corrected with Benjamini-Hochberg FDR). (D) Synergy between OsHV-1 and V. harveyi/V. rotiferianus. Left panel shows a significant increase in the mortality rate for oysters exposed to both Harveyi and OsHV-1 (in red) compared to OsHV-1 only (in black) (Kaplan-Meier survival curves, log-rank test, p = 0.0018). Right panel shows that oyster colonization by *Vibrio* is favored by OsHV-1 (Kruskal-Wallis test, p value < 0.001).

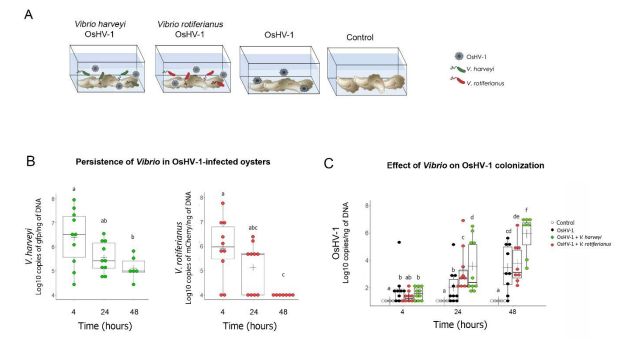


Figure 3. OsHV-1 and V. harveyi cooperate to colonize oysters

(A) Simplified mesocosm experiment using fluorescent strains of *V. harveyi* and *V. rotiferianus* (Design 2). In order to identify putative cooperative behaviors between OsHV-1 and *Vibrio*, oysters were immersed at 20°C in seawater containing OsHV-1 and fluorescent *Vibrio* (either *gfp*-labeled *V. harveyi* Th15_O_G11 or *mCherry*-labeled *V. rotiferianus* Th15_O_G05). Exposure to OsHV-1 only or immersion in seawater without introduced pathogen were used as a control. Oysters were collected at 4 h, 24 h, and 48 h, *i.e.* before mortalities occurred, to monitor pathogen load in oyster tissues. (B) Higher persistence of *V. harveyi* in OsHV-1-infected oysters. *Vibrio* loads were determined by qPCR by quantifying *gfp* and *mCherry* copies in total DNA extracted from oyster flesh in OsHV-1/*V. harveyi* and OsHV-1/*V. rotiferianus* co-infections, respectively. Each dot represents an individual oyster (t-test, p < 0.05). Only *V. harveyi* is detected at > 10⁵ copies/ng DNA over the time course of the experiment. Detection limit: 10⁴ copies/ng DNA. (C) *V. harveyi* promotes OsHV-1 replication. OsHV-1 load was measured by qPCR in the flesh of oysters exposed to OsHV-1 and *gfp*-labeled *V. harveyi* Th15_O_G11 (green) or OsHV-1 and *mCherry*-labeled *V. rotiferianus* Th15_O_G05 (red), or OsHV-1 only (black). Each dot represents an individual. (t-test, p < 0.05). A significant increase in OsHV-1 load is observed in the presence of *V. harveyi*.

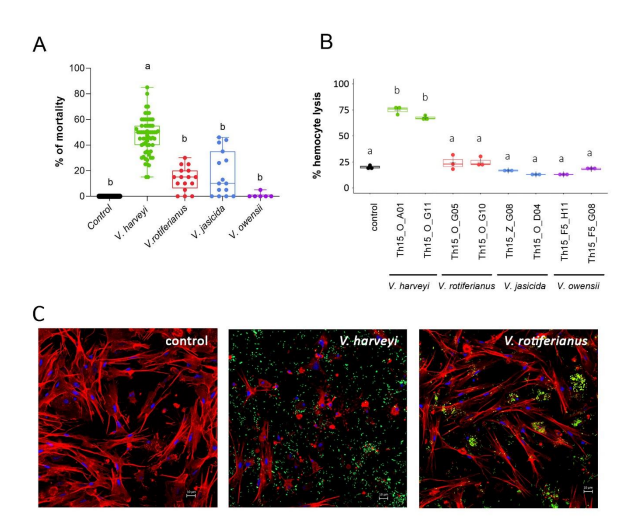


Figure 4. V. harveyi strains show high virulence potential and cytotoxicity toward oyster immune cells

(A) *V. harveyi* is significantly more virulent than other Harveyi-related species in oyster experimental infection. Oyster mortality rate (%) was measured at 24 h following an injection of Harveyi-related isolates. Mortality rates were compared for each *Vibrio* species using the Kruskal-Wallis test with post-hoc Dunn Test. Significant differences between mean values are represented by different letters (p-value < 0.0001). *V. harveyi* (n=63), *V. rotiferianus* (n=16), *V. jasicida* (n=15), *V. owensii* (n=6) (Kruskal-Wallis test, p < 0.001). *V. harveyi* isolates induced significantly higher mortalities than other strains. (B) *V. harveyi* is significantly more cytotoxic to oyster hemocytes than other species. *Vibrio* cytotoxicity was determined on monolayers of hemocytes and monitored using Sytox green labelling. Cells were incubated with bacteria at a MOI of 50:1. The maximum percentage of hemocyte lysis (%) caused by *Vibrio* is displayed. Error bars represent the standard deviation of the mean, different letters represent significant differences between means in a multiple comparison test (p<0.05, One-way ANOVA with post-hoc Tukey HSD Test). Strains of *V. harveyi* were significantly more cytotoxic than other strains. (C) *V. harveyi* but not *V. rotiferianus* causes damage to oyster hemocytes. The *Vibrio* effect on hemocytes was observed by epifluorescence microscopy. Monolayers of hemocytes were incubated with fluorescently-labeled *V. harveyi* Th15_0_G11 or *V. rotiferianus* Th15_0_G05 at a MOI of 50:1 for 2h. Actin was stained with Fluorescent-phalloidin (Red), Chromatin was stained with DAPI. *Vibrio* strains expressing fluorescent proteins are shown in Green. Cell damage was only observed with *V. harveyi*.

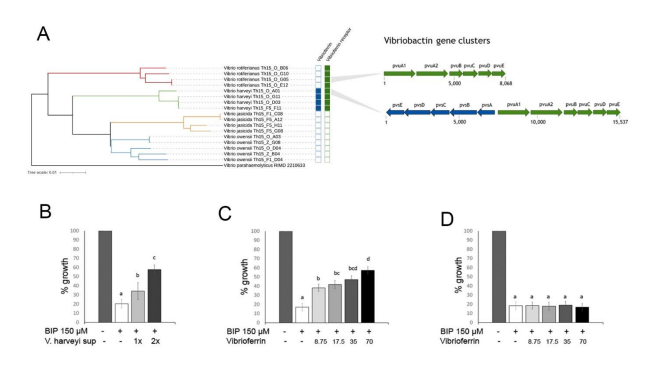


Figure 5. V. rotiferianus growth in iron-poor conditions is rescued by V. harveyi supernatant or vibrioferrin

(A) Possible cheating on vibrioferrin uptake in *V. rotiferianus*. Comparative genomics between strains of *V. harveyi* (Th15_0_G11, Th15_F5_F11, Th15_0_D03, Th15_0_A01) and *V. rotiferianus* (Th15_0_G05, Th15_0_G10, Th15_0_B06, Th15_0_E12) point to a possible cheating behavior for vibrioferrin uptake in *V. rotiferianus*, which has receptor for vibrioferrin but does not produce it. (B) *V. rotiferianus* growth in iron-depleted medium is rescued by *V. harveyi* culture supernatant. Iron-depletion was obtained by adding 150 μ M 2,2'-bipyridine (BIP) to the minimal culture medium. Dose-dependent growth rescue of strain Th15_0_G05 was achieved by adding *V. harveyi* Th15_0_G11 culture supernatant (either 1X or 2X concentrated by lyophilization). (C) *V. rotiferianus* growth in iron-depleted medium is rescued by 8.75 to 70 μ M vibrioferrin (see Fig. S8 for vibrioferrin synthesis). (D) *V. rotiferianus* growth rescue requires the iron-vibrioferrin receptor PvuA. Growth of *V. rotiferianus* Th15_0_G05 Δ pvuA1-A2 in iron-depleted medium was not rescued by addition of vibrioferrin up to 70 μ M. Letters indicate significant differences between conditions (p < 0.05, Kruskal-Wallis).

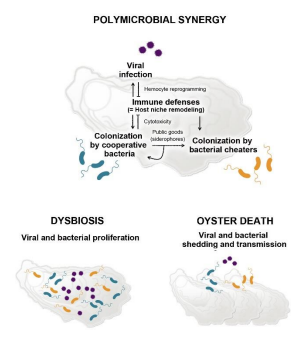


Figure 6. A systemic view of Pacific Oyster Mortality Syndrome polymicrobial synergy.

Oysters are infected by $OsHV-1 \mu var$, which impairs host immune defenses making the environment less hostile for stable bacterial colonization. Secondary bacterial colonization is enabled. The subset of cooperative bacteria that exhibit cytotoxicity toward hemocytes further dampen oyster immune defenses. This dampening is beneficial to the whole microbial community (bacteria and the OsHV-1 virus) and leads to dysbiosis. Cooperative bacteria also secrete siderophores that can be used by other members of the community who do not invest in costly mechanisms of colonization (cheaters). Cooperation between the virus and cytotoxic bacteria accelerates host death leading to the shedding of bacteria and viruses that can infect new hosts.