Factors influencing disease-induced mortality of Pacific oysters Crassostrea gigas

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ABSTRACT: Mortalities of oyster Crassostrea gigas seed associated with ostreid herpesvirus OsHV-1 µVar have been observed in many oyster-producing countries since 2008. The present study, comprised of 4 complementary experiments, aimed to identify factors associated with disease-induced oyster mortality in order to propose mitigation strategies. Our first experiment compared survival of oysters from natural spatfall with others sampled from nurseries, after thermal elevation in the laboratory from <14 to 21°C. A total of 60% of the tested wild seed batches (n = 51) were infected by OsHV-1, exhibited mortality and were able to transmit the disease to cohabited naïve oysters. Comparatively, only 1 out of the 32 tested batches sampled from nurseries presented similar characteristics. In a second experiment, we studied the effects that timing and duration of exposure to field conditions had on risk of infection and mortality in the laboratory at 21°C. Naïve oysters deployed in the field during winter and spring, when seawater temperatures were <14.7°C, showed no mortality in the laboratory, and OsHV-1 DNA was not detected by PCR. However, in oysters transferred to the field, OsHV-1 was observed when seawater temperature reached ~15.3°C. Our third experiment showed that the odds of mortality decreased with age of oysters when facing the disease. Further, we observed that odds of disease mortality decreased with water renewal and increased with the biomass of neighbouring infected oysters under controlled conditions. Based on these findings, we propose mitigation strategies in terms of the regulation of oyster movements between sites, timing of seeding and spatial planning, taking into account seawater temperature and seed origin.

KEY WORDS: Epidemiology · Risk analyses · Shellfish farming · Virus OsHV-1 µVar

INTRODUCTION

Since 2008, mass mortalities of juvenile Pacific oysters Crassostrea gigas have occurred at rearing sites along the coast of France when seawater temperature exceeds 16°C (Pernet et al. 2012, 2014b, Dégremont 2013, Petton et al. 2013). Mortality of oyster seed (<1 yr) ranges from 40 to 100%, depending on locations and batches, whereas older animals are generally much less affected (Oden et al. 2011, Pernet et al. 2012, Dégremont et al. 2013). This represents the most serious crisis the French oyster industry has faced since the mass mortality of Crassostrea angulata in the late 1960s, which lead to the introduction of C. gigas in the early 1970s (Grizel & Heral 1991). Although Pacific oyster mortalities have mainly been reported in France during this period, there have also been several cases in the UK, Australia and New Zealand (EFSA 2010, Peeler et al. 2012, Bingham et al. 2013, Jenkins et al. 2013, Paul-Pont et al. 2013, Clegg et al. 2014), and more recently in Norway (S. Mortensen pers. comm.). Results of diagnostic tests indicate that recent mortality events are associated with the detection of a particular genotype of the ostreid herpesvirus 1, named µVar (OsHV-1; Segarra et al. 2010, Jenkins et al. 2013).

Limited knowledge regarding risk factors of disease transmission and subsequent mortality has lim-
vided the development of specific protective measures for oyster farming. Additionally, unlike most other animal productions, disease curative treatments or vaccinations are not feasible in oysters (Renault 2011). As a result of the inherent dependence of the productive system on environmental hazards, oyster farmers regularly develop and conduct adaptation and risk integration strategies. Restrictions on the movements of livestock between production basins, spatial planning and density regulation in oyster beds are a few examples of farming practices implemented in response to mass mortality events (V. Le Bihan pers. comm.). In this context, additional mitigation measures are clearly needed to reduce the risks of disease transmission and mortality (Pernet et al. 2014a) in conjunction with selective breeding to improve disease resistance (Dégremont et al. 2010a).

The objective of the present study was to examine the effects of specific environmental and life-history parameters of oysters on disease susceptibility, transmission and subsequent mortality, in order to aid in mitigation strategies. To that end, 4 experiments were designed to test the effects of (1) rearing history of oyster seed, (2) timing and duration of exposure to the disease in the field, (3) age and size of oysters and (4) water renewal and biomass of infected animals on disease transmission and related mortalities in C. gigas. These experiments relied on (1) the production of standardised oyster seed under controlled conditions, (2) the use of this oyster seed for investigating the infection pressure in different environments and (3) the development of a protocol based on seawater temperature elevation in the laboratory to activate OsHV-1 in asymptomatic carriers.

The first question was addressed by investigating the effect of the origin of oysters (i.e. natural spatfall vs. nurseries) on disease transmission and mortality risk. In France, oysters originate either from natural spatfall collected along the Atlantic coast, or alternatively, from hatcheries and nurseries. In the field, oysters are unpredictably exposed to OsHV-1, whereas in hatcheries and nurseries they can be protected by means of prophylactic methods such as ultra-violet light (see details in Schikorski et al. 2011b) and seawater filtration (<5 µm, Whittington et al. 2015). Here, we tested whether the sanitary status of oyster seed is influenced by its rearing history, and explored how this could be taken into account for building disease management scenarios.

The next question we addressed was to assess the effect of timing and duration of exposure to field conditions that would result in infection and mortality in a safe environment. This parameter was evaluated in winter, spring and summer as it could potentially reflect the seasonality of virus–host interactions (EFSA 2010, Oden et al. 2011, Pernet et al. 2012, Paul-Pont et al. 2014). It is likely that during the winter when seawater temperature is far below 16°C, the risk of OsHV-1 µVar transmission to healthy oysters is low (EFSA 2010, Oden et al. 2011, Pernet et al. 2012, 2014b, Dégremont 2013, Petton et al. 2013), so that oysters could be transferred from one location to another with an acceptable risk of disease transmission. This parameter might be useful for managing the movements of livestock among shellfish culture sites, which is a common practice in France in order to optimize growth (Gouletquer & Le Moine 2002).

Thirdly, we tested the effect of age and/or size on disease susceptibility in oysters. Although several studies have reported that disease-induced mortality is lower in adults compared to other age groups (Oden et al. 2011, Peeler et al. 2012, Pernet et al. 2012, Dégremont et al. 2013, Paul-Pont et al. 2013, 2014), this generally reflects a mechanism whereby oysters that have survived a mortality event are naturally selected for resistance to that disease (Dégremont 2011, Peeler et al. 2012, Pernet et al. 2012, Dégremont et al. 2013). However, it has generally not been possible to disentangle the relative importance of age and prior OsHV-1 exposure. The effect of age and/or size on disease mortality in oysters was recently investigated by exposing healthy oysters at ages varying from 3 to 20 mo to field conditions between July 2009 and September 2011 in the Ma-rennes-Oléron Bay (France), where mortalities were occurring seasonally (Dégremont 2013). In that study, both age and/or size of oysters were negatively correlated with final cumulative mortality. In contrast, in Thau lagoon (another oyster production area in France), oysters remained highly susceptible to pathogen-related mortality pressure during their first 2 yr (Pernet et al. 2012). The novelty of our work is that oysters of different ages were exposed to the same mortality event, contrary to previous studies where the timing of exposure and age and/or size of oysters were confounded.

Finally, the last question we addressed was to study the effect of water renewal and biomass of infected animals on disease mortality of oysters. Transmission of OsHV-1 within an oyster population occurs when susceptible hosts encounter infectious particles in the environment that have been shed by neighbouring infected individuals (Schikorski et al. 2011a, Petton et al. 2013, Pernet et al. 2014a,b). We hypothesized that disease-induced mortality risk would increase with the biomass of neighbouring infected
Oysters and decrease with water renewal, reflecting concentration and dilution effects of viral particles in seawater, respectively.

**MATERIALS AND METHODS**

**Animals**

Wild individuals were collected in Fouras (Marennes-Oléron, France; 46°0’43’’ N, 1°7’3’’ W) in August 2008 and placed in mesh bags in February 2009 for transport to Paimpol (northern Brittany, France; 48°48’25’’ N, 3°0’23’’ W) until February 2010. The animals were then moved to a grow-out farm located at Aber Benoit, (northern Brittany, France; 48°34’30’’ N, 4°36’18’’ W) until being used as broodstock. These animals were exposed to the disease during the spring 2009 and suffered ca. 75% mortality (Petton et al. 2013).

Between March 2010 and April 2011, 10 batches of 40 adult oysters each were transferred every 6 wk to the Ifremer facilities in Argenton (Brittany, France; 48°31’16’’ N, 4°46’2’’ W) for a conditioning period of 32 to 45 d (Table 1). These animals were held in 500 l flow-through tanks with seawater held at a constant temperature of 17°C and enriched with a phytoplankton mixture. Seawater was treated with UV and filtered through 1 µm mesh. The daily mixed diet consisted of *Isochrysis affinis galbana* (CCAP 927/14) and *Chaetoceros gracilis* (UTEX LB2658) 1:1 in dry weight. Once the oysters were reproductively mature, gametes from 30 individuals (1/3 males, 2/3 females), obtained by stripping, were mixed in a 5 l jar at 50 spermatozoaids per oocyte. The fertilized oocytes completed their embryonic development in 150 l tanks filled with 1 µm filtered and UV-treated seawater at 21°C for 24 h. The D-larvae were then collected and reared in flow-through rearing systems at 25°C (Rico-Villa et al. 2008). At the end of the pelagic phase (12 d), competent larvae were collected on a 225 µm sieve and allowed to settle on cultch. Post-larvae were maintained in downwelling systems where they were continuously supplied with enriched seawater. After 5 d, the cultchless seed were collected on 400 µm mesh and reared at 25°C depending on the experiment. In the larval and post-larval stages, the oysters were fed the same diet as the broodstock at a concentration of 1500 µm³ µl⁻¹ (Rico-Villa et al. 2009). Overall, 10 batches of standardized oyster seed (3 to 6 mo old, 0.2 to 1.2 g wet mass) were produced every 6 wk for use in the experiments. They all remained free of any abnormal mortality, OsHV-1 DNA was not detected and the relative concentration of vibrios was low (Table 1). Thus, the 10 standardised batches were considered naïve with regard to OsHV-1 µVar. The likely minimum level of disease prevalence was <30% (with 95% confidence level), considering that 10 oysters were analysed for OsHV-1 DNA detection by qPCR, and assuming that there is no loss of sensitivity due to pooling and that the population of oysters is infinite (Pfeiffer 2010).

<table>
<thead>
<tr>
<th>Batch no.</th>
<th>Conditioning</th>
<th>Fertilization</th>
<th>Experiments</th>
<th>Type</th>
<th>Locations</th>
<th>Sites</th>
<th>Temperature (°C)</th>
<th>Age (mo)</th>
<th>Mass (mg ind.⁻¹)</th>
<th>Oysters</th>
<th>OsHV-1 DNA detection</th>
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<tr>
<td>1</td>
<td>23/03/10</td>
<td>03/05/10</td>
<td>15/07/10</td>
<td>A</td>
<td>Arg</td>
<td>Arg</td>
<td>21.0</td>
<td>2.4</td>
<td>310 ± 10</td>
<td>nd</td>
<td>1.34 ± 0.50</td>
</tr>
<tr>
<td>2</td>
<td>28/04/10</td>
<td>09/06/10</td>
<td>25/08/10</td>
<td>A, C</td>
<td>Arg, AB</td>
<td>Arg</td>
<td>21.0 na</td>
<td>2.5</td>
<td>260 ± 10</td>
<td>nd</td>
<td>1.67 ± 0.40</td>
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<tr>
<td>3</td>
<td>07/06/10</td>
<td>19/07/10</td>
<td>08/10/10</td>
<td>A, C</td>
<td>Arg, AB, BB</td>
<td>Arg</td>
<td>21.0 na 16.3</td>
<td>2.7</td>
<td>337 ± 12</td>
<td>nd</td>
<td>0.25 ± 0.10</td>
</tr>
<tr>
<td>4</td>
<td>17/07/10</td>
<td>30/08/10</td>
<td>22/11/10</td>
<td>A, B, C</td>
<td>Arg, AB, BB</td>
<td>Arg</td>
<td>21.0 na 11.6</td>
<td>2.8</td>
<td>263 ± 12</td>
<td>nd</td>
<td>0.23 ± 0.10</td>
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<tr>
<td>5</td>
<td>30/08/10</td>
<td>11/10/10</td>
<td>03/01/11</td>
<td>A, C</td>
<td>Arg, AB, BB</td>
<td>Arg</td>
<td>21.0 9.8 8.1</td>
<td>2.8</td>
<td>273 ± 6</td>
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<td>0.04 ± 0.20</td>
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<td>17/02/11</td>
<td>A, C</td>
<td>Arg, AB, BB</td>
<td>Arg</td>
<td>21.0 9.4 8.7</td>
<td>2.9</td>
<td>297 ± 12</td>
<td>nd</td>
<td>0.31 ± 0.00</td>
</tr>
<tr>
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<td>22/11/10</td>
<td>05/01/11</td>
<td>05/04/11</td>
<td>A, C</td>
<td>Arg, AB, BB</td>
<td>Arg</td>
<td>21.0 11.5 12.9</td>
<td>3.0</td>
<td>333 ± 15</td>
<td>nd</td>
<td>0.11 ± 0.10</td>
</tr>
<tr>
<td>8</td>
<td>10/01/11</td>
<td>22/02/11</td>
<td>13/05/11</td>
<td>A</td>
<td>Arg</td>
<td>Arg</td>
<td>21.0 14.6 16.4</td>
<td>2.6</td>
<td>343 ± 6</td>
<td>nd</td>
<td>0.00 ± 0.00</td>
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<td>2.7</td>
<td>353 ± 6</td>
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<td>0.32 ± 0.10</td>
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<td>A</td>
<td>Arg</td>
<td>Arg</td>
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<td>3.0</td>
<td>327 ± 6</td>
<td>nd</td>
<td>1.08 ± 0.70</td>
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</table>

Table 1. Characteristics of oyster seed used in Expts A–D. OsHV-1 DNA detection was performed on 2 pools of 5 oysters; only results from unexposed oysters are presented here. Bacterial analyses were conducted on 3 to 8 pooled oysters in duplicate or in triplicate. Biomass and *Vibrio* spp. data are means ± SD. Arg: Argenton; AB: Aber Benoit; BB: Bay of Brest; nd: not detected; na: not available. Age was measured as months from fertilization to onset of experiment. Dates are presented as dd/mm/yy.

*aOne replicate only, other data missing*
Experimental design

Expt A: effect of oyster seed origin on disease transmission and mortality risk

The objective of Expt A was to investigate whether disease transmission and the related mortality of oysters is influenced by their rearing history (i.e. exposure in the field versus rearing under hatchery/nursery conditions) (Fig. 1). To test this hypothesis, 83 batches of seed (<1 yr) were collected between 2010 and 2014 (see Supplement 1 at www.int-res.com/articles/supp/q006p205_supp.pdf). A total of 51 batches were sampled from natural collection sites in France: Thau lagoon (3), Arcachon (10), Marennes-Oléron (14), Bourgneuf (7), Vilaine estuary (4), Quiberon (1) and the Bay of Brest (12). An additional 32 batches were obtained from nurseries—either public research facilities: Ifremer Argenton (24), Ifremer La Tremblade (2), or commercial hatcheries (6). Wild oyster seed were collected during the winter or early spring when seawater temperature was below 14°C; no abnormal mortality occurred. Oyster seed from nurseries were collected all year, reflecting the fact that hatchery production of oysters is usually not synchronised with natural spawning. Oyster seed sampled in nurseries were 3 mo old, and their wet weight varied from 0.2 to 1.2 g.

Animals were directly transferred to the Ifremer facilities in Argenton, where they were placed at 21°C in order to re-activate the virus in infected individuals. Survival was followed for 30 d. Note that oysters originating from Argenton were held at 21°C all year, so they were previously acclimated for 10 d at 13°C before being re-exposed at 21°C.

Additionally, 38 batches tested in 2013 and 2014 were placed in cohabitation with naïve oysters to further investigate whether transmission of the disease occurred (see Table S1 in the Supplement). These cohabitation trials were designed based on previous results which showed that it is possible to transmit disease and related mortality to healthy Crassostrea gigas seed in cohabitation experiments, by using oysters infected by a brief exposure to field conditions in which mortalities were occurring (Petton et al. 2013). The tested oyster seed (n = 150) were placed on one side of each tank and naïve oysters (n = 100) were placed downstream, on the opposite side. These naïve oysters were hereafter referred to as ‘challenged’ animals.

Live and dead oysters were counted at the start of the experiment and after 30 d. Dead animals were removed from the tank at each count. Duplicate pools of 5 oysters from each batch were sampled for OsHV-1 DNA detection (1) at the time of collection and (2) when mortality occurred, or after 10 to 15 d of acclimation at 21°C.

Expt B: determination of the timing and duration of field exposure required to become diseased

The objective of this experiment was to determine the timing and duration of exposure to field conditions (where disease mortality of oysters naturally occurs) that is required for otherwise healthy individuals to become infected and exhibit mortality in a safe and controlled environment (Fig. 1). Sub-samples of oysters originating from batches 4, 8 and 9 were transferred on 22 November 2010, 31 May 2011 and 4 July 2011 respectively (Table 1) to a farming area located in the Bay of Brest at Pointe du Château (48° 20’ 6″ N, 4° 19’ 6″ W). Experimental oysters were placed in duplicate mesh bags (ca. 150 ind. bag⁻¹) attached to iron tables. Seawater temperatures were 11.6, 16.1 and 19.2°C, respectively (Table 1). The experimental oysters were exposed to field conditions for 42, 73, 102, 135, 149 or 162 d (batch 4); 6, 9, 13, 17 or 20 d (batch 8); or 4, 6, 8, 10 or 12 d (batch 9). The field-exposed oysters did not show significant mortality during the exposure periods. Sub-samples of the exposed oyster batches were left in the field to examine whether they suffered mortalities later.

Following field exposure, oysters were transferred back to the Ifremer facilities in Argenton where they were placed in rectangular 23 l plastic tanks at 21°C. Each tank contained all oysters originating from 1 mesh bag (n = 2 tanks for each condition with 150 ind. tank⁻¹). Each tank was supplied with 1 µm filtered seawater exposed to UV irradiation flowing at 100 ml min⁻¹ (Petton et al. 2013). Live and dead animals were counted every 1 to 2 d, and dead animals were removed from the tanks. For each batch, survival of unexposed oysters was followed in duplicate tanks (controls). OsHV-1 DNA detection was conducted on 5 pooled oysters sampled at the end of field exposure (the onset of the laboratory monitoring) for each replicate (n = 2). Both field-exposed and control oysters were analysed. Additionally, oysters that were left in the field for the entire duration of the experiment were sampled and analysed for OsHV-1 DNA detection by PCR (see ‘Quantification of OsHV-1 DNA’ below) when mortality occurred.
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Expt A: Effect of origin of oyster seed on disease mortality risk

Exposure at 21°C in the lab in cohabitation with naïve oysters
Sampling for OsHV-1 DNA detection
Survival monitoring (30d)
Sampling for OsHV-1 DNA detection

Naïve oysters
Start of field exposure

Batch 4
22/11/10
149
192
139
122
113
77
42

Batch 8
31/05/11
12
11
10
9

Batch 9
4/7/11
12
11
10
8
6

Expt B: Determination of the timing and duration of field exposure required to become diseased

Oyster seed collection
(Wild (n=51)
Nursery (n=32)

Expt C: Effect of age and/or size of oyster seed on disease susceptibility of oysters

Date of transfer in the field
Sampling times for survival

Expt D: Effect of water renewal and biomass of infected oysters on disease transmission and related mortalities

Water renewal

Fig. 1. Schematic of designs used in Expts A–D to identify factors associated with disease-induced mortality in Pacific oysters Crassostrea gigas, in particular in the presence of ostreid herpesvirus OsHV-1 µVar
Expt C: effect of age and/or size of oyster seed on disease susceptibility

Sub-samples of 3 mo old oysters originating from batches 2 to 7 were transferred to the Bay of Brest at Pointe du Château, and to Aber Benoit on 25 August, 8 October and 22 November 2010, and on 3 January, 17 February and 5 April 2011 (Table 1). Oysters were placed in duplicate mesh bags (ca. 150 ind. bag⁻¹) attached to iron tables. These oysters did not show significant mortality between deployment and May 2011, except for animals deployed at the Bay of Brest on 25 August 2010 (batch 2), which exhibited 75% mortality at the beginning of September. This batch was therefore not included in Expt C at the Bay of Brest. The age of the oysters was ca. 4.3, 5.8, 7.1, 8.5, 9.9 and 11.2 mo.

Survival of oysters was recorded on 2 and 17 May 2011 at the Bay of Brest and Aber Benoît respectively, when the seawater temperature reached 16°C (a threshold temperature above which disease transmission is optimal and mortalities occur) and at the onset of the mass mortality (18 May and 15 June). Subsequently, survival of oysters was followed on 5 July, 4 August and 13–30 September 2011 in Aber Benoît and Bay of Brest, and every 4 to 6 mo thereafter until 4–5 February 2014. To evaluate the effect of size at age on survival, shell lengths of live and dead oysters contained in 1 of the 2 bags transferred to Aber Benoît were measured on 15 June 2011, when mass mortality occurred.

In this experiment, age was confounded with duration of field exposure before the onset of mortality. Therefore, to separate these 2 effects, a complementary experiment was performed (see Supplement 2 at www.int-res.com/articles/suppl/q006p205_supp.pdf). Briefly, oysters from batches 5 and 7 (aged 7.1 and 4.3 mo respectively at the onset of mortality) were transferred twice to the Bay of Brest: oysters from batch 5 were transferred on 3 January (long exposure = 89 d) and later on 5 April (short exposure = 41 d), whereas those from batch 7 were transferred on 5 April (long exposure = 41 d) and later on 3 May (short exposure = 13 d). Survival of these animals was followed for 1 yr until 3 May 2012.

Expt D: effect of water renewal and biomass of infected oysters on disease transmission and mortality

The objective of Expt D was to investigate the effect of water renewal and biomass of infected animals on disease transmission and related mortalities of oysters under experimental tank conditions. Subsamples of oysters originating from batch 10 were exposed to field conditions in the Bay of Brest at Pointe du Château on 12 September and 1 October 2011 for the water renewal and biomass experiment respectively, when seawater temperature was 18.4 and 16.1°C, respectively.

On 10 October, these previously field-exposed oysters were transferred back to the Ifremer facilities in Argenton where they were placed randomly in duplicate rectangular 23 l plastic tanks in contact with naïve oysters at 3 different water renewal rates (48, 120 and 280 ml min⁻¹). A total of 100 exposed oysters (total biomass = 160 g) were placed on one side of each tank, whereas another 135 naïve oysters (160 g) were placed downstream, on the opposite side.

Similarly, on 28 October, 3 different biomasses of exposed oysters (50, 100 and 200 g) were placed in duplicate 23 l tanks in contact with naïve oysters (75 animals, total biomass = 125 g, hereafter referred to as ‘laboratory challenged’ animals). Water renewal was maintained at a constant rate of 100 ml min⁻¹. Live and dead oysters were counted at the start of the experiments and after 1, 2, 4, 7, 11, 14 and 19 d (for the water renewal experiment) or after 7, 11 and 19 d (for the biomass experiment). Dead animals were removed from the tank at each count. Exposed oysters were sampled for OsHV-1 DNA detection at the onset of the experiment and after 2 and 4 d.

Monitoring of seawater temperature in the field

Seawater temperatures were provided by the Resco/Velyger Ifremer networks (http://www.ifremer.fr/velyger). Temperatures have been recorded every 30 min since March 2009 using multi-parameters probes (NKE Instrumentation) deployed at our experimental sites in the Bay of Brest at Pointe du Château (48°20’06.19″N, 4°19’06.37″W) and at Aber Benoît (48°34’29.976″N, 4°36’18.378″W).

OsHV-1 and vibrios analyses

Oysters sampled for OsHV-1 DNA detection were stored at ~20°C until analysis. DNA extraction and OsHV-1 detection by PCR were performed by IDHESA Bretagne Océane (Quimper).

Quantification of OsHV-1 DNA

Total DNA was extracted from an aliquot of tissue sample using a QIAgen QIAaamp tissue mini kit, according to the manufacturer’s protocol. The extracted
DNA was stored at −20°C prior to analysis. The detection and quantification of OsHV-1 DNA was carried out using a previously published real-time PCR protocol (Pepin et al. 2008). Briefly, this protocol uses SYBR® Green chemistry with specific DPFor/Rev primers targeting the region of the OsHV-1 genome predicted to encode a DNA polymerase catalytic subunit (Webb et al. 2007). The specificity and sensitivity of the detection test using these primers is similar to those reported by Pepin et al. (2008) (T. Renault pers. comm.). The method used in our study was the recommended method for reasons of availability, utility, and diagnostic specificity and sensitivity for OsHV-1 detection (OIE 2012, www.oie.int/fileadmin/Home/eng/Health_standards/aahm/2010/2.4.09_INF_OSTREID_HERPES.pdf). Results were expressed as viral DNA copy numbers mg−1 wet tissue.

Targeted detection of OsHV-1 µVar

The standard real-time PCR using SYBR Green cannot differentiate OsHV-1 reference from OsHV-1 µVar. Therefore, specific primers were used to distinguish the ‘reference’ and ‘µVar’ genotypes by comparison with positive controls (see Appendix B in EFSA 2010, Council regulation 175/2010). These complementary OsHV-1 µVar specific PCR analyses were performed every 30 samples in which OsHV-1 DNA had first been detected. The OsHV-1 µVar-specific PCR analyses detected the OsHV-1 µVar genotype only, which agrees well with the fact that in 2009 the OsHV-1 µVar had fully replaced the reference OsHV-1 genotype in seed presenting mortality at all French oyster production sites (Segarra et al. 2010, Martenot et al. 2011, Renault et al. 2012). Therefore, in the present study we refer to ‘OsHV-1’ as ‘OsHV-1 µVar’.

Quantification of vibrios

Whole tissue samples of a pool of 3 to 8 oysters per batch, diluted in sterilized seawater (1:100), were spread on marine broth medium and thiosulphate citrate bile salts-sucrose (TCBS) in Petri dishes to quantify cultivable bacteria, and more specifically, vibrios (Petton et al. 2013). The bacteriological media used in this study were obtained from Difco Laboratories. The plates were incubated at 22.5°C for 6 d in marine broth, and for 2 d in TCBS before counting the number of colony forming units (CFU, Table 1). Data are expressed as % Vibrio spp. (100 × CFU of vibrios / CFU of total cultivable bacteria).

Statistical analyses

Statistical analyses were conducted using LIFETEST, LOGISTIC and PHREG procedures of the SAS software package (SAS v.9.4, SAS institute). A significance threshold of \( \alpha = 0.05 \) was adopted for all statistical tests.

Lifetest

Nonparametric estimates of the survivor function were computed by the Kaplan-Meier method (Kaplan & Meier 1958). Survival time was measured as days from the onset of the experiment, when oysters were transferred back to the Ifremer facilities in Argenton at 21°C. The data were read as the number of dead animals within each experimental unit at each time interval. Survival curves of oysters were compared among treatments (Expt B: duration of field exposure at each season; Expt C: age and/or size, duration of field exposure; Expt D: water renewal and biomass of infected oysters) and status (Expt D: previously field-exposed or laboratory-challenged). The resulting survival estimates were compared by using the log-rank test of homogeneity of strata.

Logistic regression

A logistic regression model was used to investigate the effect of exposure to field conditions on the occurrence of mortality, and the effect of detecting OsHV-1 DNA in oyster tissues before and during exposure at 21°C in the laboratory (Expt A). The model predicts the probability of a 1 for mortality and a 0 for absence of mortality. Mortality was considered significant when >5%. The variable origin was coded 0 if oysters were from a nursery, 1 for wild oysters. OsHV-1 DNA detection in oysters was coded 0 if not detected, 1 for detected.

Cox regression

The survival time curves of oysters were compared using the Cox regression model (Cox 1972), after adjustment for the effect of some static covariates such as age for each site (Expt C) or water renewal and biomass of infected oysters and status (previously field-exposed or laboratory-challenged, Expt D). The proportionality of hazards (PH) was checked with martingale residuals (Lin et al. 1993, Lee &
Wang 2013). Because the PH assumption was violated, time-dependent covariates representing the interaction of the original covariates and time were added to the model. Time (t) was defined as dichotomous: \( t \leq 136 \text{ d} \) or \( t > 136 \text{ d} \) (Expt C) and \( t \leq 7 \text{ d} \) or \( t > 7 \text{ d} \) (Expt D). These time intervals contained a sufficient number of deaths to estimate regression coefficients reliably (Anderson & Senthilselvan 1982, Hess 1995). Custom hazard ratios were produced by means of polynomial (linear) contrasts.

### RESULTS

#### Effect of origin of oyster seed on disease mortality risk (Expt A)

The survival of oysters collected between 2010 and 2014 from 7 seed collection sites (51 batches) and 5 nurseries (32 batches), when seawater temperature was <14°C, was followed for 30 d at 21°C under laboratory conditions in duplicate tanks (i.e. trials). OsHV-1 DNA detection analyses were conducted twice: at the time of collection and when mortality occurred, or after 10 to 15 d of acclimation at 21°C. Most batches collected from nurseries did not exhibit mortality at 21°C under laboratory conditions, and OsHV-1 DNA was not detected during survival monitoring. Furthermore, challenged oysters placed in cohabitation for 30 d with oysters collected from hatcheries showed no mortality; thus there was no disease transmission. Only 1 batch (no. 10) out of 32 tested (2/63 trials; Table 2) sampled from a nursery showed significant mortality and was positive for the detection of OsHV-1 DNA (see Supplement 1).

By contrast, 31/51 batches collected from the wild (53/101 trials) showed significant mortalities at 21°C in at least 1 trial (Table 2). These mortalities corresponded to the detection of OsHV-1 DNA in oyster tissues before and/or during survival monitoring in 28/31 batches (50/53 trials; Table 2). In these cases, survival varied from 18 to 95% with a mean (± SD) of 54 ± 23%. Also, challenged oysters placed in cohabitation with diseased wild oysters showed significant mortalities in 28/31 trials, suggesting disease transmission occurred. It is noteworthy that mortalities were associated with the detection of OsHV-1 DNA in only 3/53 trials. However, only one of these trials showed high mortality in both collected and challenged oysters (survival was 60.2% in collected oysters and 75.2% in challenged animals in 1 of 2 trials of batch 68; see Supplement 1). The other 2 trials exhibited 91.3 and 94.5% survival, and challenged oysters showed no mortalities (see batches 69 and 66, respectively in Supplement 1).

Lack of significant mortality in wild oysters at 21°C was associated with the absence of OsHV-1 DNA in their tissues in 28/48 trials, and the challenged oysters placed in cohabitation with these wild oysters showed no mortality (Table 2). Conversely, OsHV-1 DNA had been detected in wild oysters although no mortality occurred in 20/48 trials, and the challenged oysters showed no mortality, suggesting there was no disease transmission.

Overall, the odds of mortality at 21°C under laboratory conditions were 27 times higher in wild-collected oysters compared to those from nurseries (Table 3). Detection of OsHV-1 DNA in oysters at the time of collection was not predictive of mortality risk. When oysters were exposed at 21°C, odds of mortality were 66 times higher when OsHV-1 DNA was detected than when it was not.

<table>
<thead>
<tr>
<th>Origin</th>
<th>Mortality</th>
<th>OsHV-1 DNA detection</th>
<th>No. of trials</th>
<th>Mortality in challenged oysters</th>
</tr>
</thead>
<tbody>
<tr>
<td>Nursery</td>
<td>No</td>
<td>No</td>
<td>61 (37.2)</td>
<td>50 (31.1)</td>
</tr>
<tr>
<td></td>
<td>Yes</td>
<td>Yes</td>
<td>2 (1.2)</td>
<td>2 (1.2)</td>
</tr>
<tr>
<td>Wild</td>
<td>No</td>
<td>No</td>
<td>28 (17.1)</td>
<td>4 (2.4)</td>
</tr>
<tr>
<td></td>
<td>No</td>
<td>Yes</td>
<td>15 (9.1)</td>
<td>9 (5.5)</td>
</tr>
<tr>
<td></td>
<td>Yes</td>
<td>No</td>
<td>1 (0.6)</td>
<td>0 (0.0)</td>
</tr>
<tr>
<td></td>
<td>Yes</td>
<td>Yes</td>
<td>4 (2.4)</td>
<td>3 (1.8)</td>
</tr>
<tr>
<td></td>
<td>Yes</td>
<td>No</td>
<td>3 (1.8)</td>
<td>1 (0.6)</td>
</tr>
<tr>
<td></td>
<td>Yes</td>
<td>Yes</td>
<td>39 (23.8)</td>
<td>17 (10.4)</td>
</tr>
<tr>
<td></td>
<td>Yes</td>
<td>Yes</td>
<td>11 (6.7)</td>
<td>2 (1.2)</td>
</tr>
<tr>
<td>Total</td>
<td>–</td>
<td>–</td>
<td>164 (100.0)</td>
<td>47 (28.7)</td>
</tr>
</tbody>
</table>

---

Table 2. Expt A. Mortality at 21°C and detection of ostreid herpesvirus 1 (OsHV-1) DNA in Pacific oyster *Crassostrea gigas* seed collected in the wild (51 batches) and in nurseries (32 batches). Data are number of trials corresponding to each combination of origin, mortality and OsHV-1 DNA detection. The numbers in parentheses indicate percentages based on the total amount of trials. Two trials were generally conducted for each batch of oysters (n = 83 batches, 164 trials; see Supplement 1 at www.int-res.com/articles/supp/q006p205_supp.pdf for further details). Mortality was also recorded in challenged oysters placed in cohabitation with a sub-sample of oysters collected in the wild (33 batches) and in nurseries (5 batches) to investigate whether transmission of the disease occurred. Mortality was considered significant when >5%. Values in **bold** indicate the most common combinations (>10%); nd: not determined.
The oysters deployed in the Bay of Brest at Pointe du Chateau on 22 November 2010 for 42 to 149 d (until 20 April 2011 when average seawater temperature was 14.8°C) exhibited 100% survival when exposed to a temperature elevation at 21°C in the laboratory (Fig. 2A). OsHV-1 DNA was not detected in these animals (0 out of 10 pools of 5 oysters; Table 4). Survival of oysters left in the field for 162 d (until 3 May 2011, average seawater temperature was 15.3°C) and then transferred to the laboratory at 21°C was only 28.4% in one replicate but 100% in the other (mean = 64.2%; Fig. 2A). The level of OsHV-1 DNA in oysters sampled in the mesh bag where mortality occurred was $4.1 \times 10^6$ copies mg$^{-1}$ wet tissue (Table 4), which is 2 orders of magnitude higher than the threshold value of $10^4$ DNA copies mg$^{-1}$ wet tissue, at which OsHV-1 is considered to be involved in mortality (Oden et al. 2011, Schikorski et al. 2011a). It is noteworthy that oysters left in the field exhibited mortality was 14.8°C) exhibited 100% survival when exposed to a temperature elevation at 21°C in the laboratory (Fig. 2A). OsHV-1 DNA was not detected in these animals (0 out of 10 pools of 5 oysters; Table 4). Survival of oysters left in the field for 162 d (until 3 May 2011, average seawater temperature was 15.3°C) and then transferred to the laboratory at 21°C was only 28.4% in one replicate but 100% in the other (mean = 64.2%; Fig. 2A). The level of OsHV-1 DNA in oysters sampled in the mesh bag where mortality occurred was $4.1 \times 10^6$ copies mg$^{-1}$ wet tissue (Table 4), which is 2 orders of magnitude higher than the threshold value of $10^4$ DNA copies mg$^{-1}$ wet tissue, at which OsHV-1 is considered to be involved in mortality (Oden et al. 2011, Schikorski et al. 2011a). It is noteworthy that oysters left in the field exhibited

<table>
<thead>
<tr>
<th>Source of variation</th>
<th>df</th>
<th>Estimate</th>
<th>SE</th>
<th>$\chi^2$</th>
<th>p</th>
<th>Odds ratio</th>
<th>95% CI</th>
</tr>
</thead>
<tbody>
<tr>
<td>Origin of oysters</td>
<td></td>
<td>−3.3</td>
<td>0.7</td>
<td>24.3</td>
<td>&lt;0.001</td>
<td>27.1</td>
<td>7.2–102.9</td>
</tr>
<tr>
<td>OsHV-1 initial</td>
<td></td>
<td>0.2</td>
<td>0.6</td>
<td>0.1</td>
<td>0.809</td>
<td>1.2</td>
<td>0.3–4.0</td>
</tr>
<tr>
<td>OsHV-1 at 21°C</td>
<td></td>
<td>4.2</td>
<td>0.6</td>
<td>45.2</td>
<td>&lt;0.001</td>
<td>66.1</td>
<td>19.5–224.1</td>
</tr>
</tbody>
</table>

Table 3. Expt A. Logistic regressions examining the effects on survival of the origin of Pacific oyster *Crassostrea gigas* seed (wild or nursery) and detection of OsHV-1 DNA before and during thermal elevation in the laboratory at 21°C. For each factor, the following elements are given: its parameter estimate, standard error, chi-square and the resulting p-value for the Type II test from the complete model. The corresponding instantaneous odds ratio and the confidence interval are also provided. Significant p-values (p < 0.05) are in bold

**Determination of the timing and duration of field exposure required to become diseased (Expt B)**

The oysters deployed in the Bay of Brest at Pointe du Chateau on 22 November 2010 for 42 to 149 d (until 20 April 2011 when average seawater temperature was 14.8°C) exhibited 100% survival when exposed to a temperature elevation at 21°C in the laboratory (Fig. 2A). OsHV-1 DNA was not detected in these animals (0 out of 10 pools of 5 oysters; Table 4). Survival of oysters left in the field for 162 d (until 3 May 2011, average seawater temperature was 15.3°C) and then transferred to the laboratory at 21°C was only 28.4% in one replicate but 100% in the other (mean = 64.2%; Fig. 2A). The level of OsHV-1 DNA in oysters sampled in the mesh bag where mortality occurred was $4.1 \times 10^6$ copies mg$^{-1}$ wet tissue (Table 4), which is 2 orders of magnitude higher than the threshold value of $10^4$ DNA copies mg$^{-1}$ wet tissue, at which OsHV-1 is considered to be involved in mortality (Oden et al. 2011, Schikorski et al. 2011a). It is noteworthy that oysters left in the field exhibited

<table>
<thead>
<tr>
<th>Source of variation</th>
<th>df</th>
<th>Estimate</th>
<th>SE</th>
<th>$\chi^2$</th>
<th>p</th>
<th>Odds ratio</th>
<th>95% CI</th>
</tr>
</thead>
<tbody>
<tr>
<td>Origin of oysters</td>
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<td>−3.3</td>
<td>0.5</td>
<td>35.3</td>
<td>&lt;0.001</td>
<td>27.1</td>
<td>7.2–102.9</td>
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<tr>
<td>OsHV-1 initial</td>
<td></td>
<td>0.2</td>
<td>0.6</td>
<td>0.1</td>
<td>0.809</td>
<td>1.2</td>
<td>0.3–4.0</td>
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<tr>
<td>OsHV-1 at 21°C</td>
<td></td>
<td>4.2</td>
<td>0.6</td>
<td>45.2</td>
<td>&lt;0.001</td>
<td>66.1</td>
<td>19.5–224.1</td>
</tr>
</tbody>
</table>

Table 3. Expt A. Logistic regressions examining the effects on survival of the origin of Pacific oyster *Crassostrea gigas* seed (wild or nursery) and detection of OsHV-1 DNA before and during thermal elevation in the laboratory at 21°C. For each factor, the following elements are given: its parameter estimate, standard error, chi-square and the resulting p-value for the Type II test from the complete model. The corresponding instantaneous odds ratio and the confidence interval are also provided. Significant p-values (p < 0.05) are in bold

**Determinant of the timing and duration of field exposure required to become diseased (Expt B)**

The oysters deployed in the Bay of Brest at Pointe du Chateau on 22 November 2010 for 42 to 149 d (until 20 April 2011 when average seawater temperature was 14.8°C) exhibited 100% survival when exposed to a temperature elevation at 21°C in the laboratory (Fig. 2A). OsHV-1 DNA was not detected in these animals (0 out of 10 pools of 5 oysters; Table 4). Survival of oysters left in the field for 162 d (until 3 May 2011, average seawater temperature was 15.3°C) and then transferred to the laboratory at 21°C was only 28.4% in one replicate but 100% in the other (mean = 64.2%; Fig. 2A). The level of OsHV-1 DNA in oysters sampled in the mesh bag where mortality occurred was $4.1 \times 10^6$ copies mg$^{-1}$ wet tissue (Table 4), which is 2 orders of magnitude higher than the threshold value of $10^4$ DNA copies mg$^{-1}$ wet tissue, at which OsHV-1 is considered to be involved in mortality (Oden et al. 2011, Schikorski et al. 2011a). It is noteworthy that oysters left in the field exhibited

**Fig. 2. Expt B. Survival of Pacific oysters *Crassostrea gigas* under laboratory conditions at 21°C according to the timing (A: 22 November 2010, B: 31 May 2011, C: 4 July 2011) and the duration (4 to 162 d) of exposure in the Bay of Brest. Survival time was measured as days from the end of the field exposure period, i.e. the onset of the temperature elevation to 21°C**
70% mortality on 16 May, concomitant with high levels of OsHV-1 DNA (data not shown).

The oysters deployed at the Bay of Brest on 31 May for only 6 to 9 d while seawater temperature was >16°C exhibited 100% survival when exposed to a temperature increase to 21°C in the laboratory (Fig. 2B), and OsHV-1 DNA was not detected in their tissues (Table 4). By contrast, mean survival of oysters left in the field for 13 to 20 d was only 24.0% in the laboratory at 21°C, and OsHV-1 DNA was detected in all oyster samples (Fig. 2B, Table 4). Overall, final survival of oysters decreased with duration of field exposure. Oysters left in the field exhibited high levels of OsHV-1 DNA in their tissues after 13 d of exposure (1.7 ± 3.3 × 10^6 copies mg⁻¹ wet tissue), and survival decreased markedly thereafter (54 ± 2% after 21 d) (data not shown).

### Effect of age and size of oyster seed on disease susceptibility (Expt C)

Survival of oysters of 4.3 to 11.2 mo of age was followed from May 2011 until February 2014 in Aber Benoît and in the Bay of Brest. In the Bay of Brest, daily average seawater temperature fluctuated seasonally between 7.5 and 21.0°C, whereas in Aber Benoît it varied from 6.5 to 17.0°C (Fig. 3).

Final survival of oysters maintained at Aber Benoît was 44.1% compared to 25.7% in the Bay of Brest (log-rank, p < 0.001). Also, the onset of oyster mortality occurred between 2 and 17 May at the Bay of Brest, where daily average maximum seawater temperature was 17.2°C, whereas in Aber Benoît mortality began later (between 17 May and 15 June), while daily average maximum seawater temperature was only 15.0°C (Fig. 3). The onset of mortality coincided with the detection of OsHV-1 DNA in oyster tissues both in the Bay of Brest (see results of Expt A) and in Aber Benoît, where oysters exhibited 2.1 × 10^7 ± 2.3 × 10^7 copies OsHV-1 DNA mg⁻¹ tissue (n = 2 samples).

Survival of oysters varied as a function of age and time both at Aber Benoît (Cox non-proportional hazard model, age × time [>136 d], Wald χ² = 114.0, p < 0.001) and Bay of Brest (age × time [>136 d], Wald χ² = 54.5, p < 0.001). In the first 136 d after disease outbreak, parameter estimates for linear effects of age on survival were significant at both sites (Table 5). Based on these estimates, we calculated that an increase of 5.7 mo of age in oysters reduced the odds of mortality by 44% (7.7% mo⁻¹) in Aber Benoît. Similarly, in the Bay of Brest, odds of mortality during the first 136 d after disease outbreak decreased by 35% with an age increase of 4.4 mo (7.9% mo⁻¹).

During the next time period (136 to 995 d), parameter estimates for a linear effect of age were either not significant (Aber Benoît) or only marginally significant (Bay of Brest; Table 5), highlighting the fact that the effect of age becomes progressively less important with time. In Bay of Brest, the probability of mortality increased with the age of oysters, reflecting the fact that it was higher in 9.9 mo old oysters compared to that of the other age groups (Fig. 3, Table 5).

### Table 4. Expt B. Detection of OsHV-1 DNA in Pacific oysters *Crassostrea gigas* exposed to field conditions in the Bay of Brest on 22 November 2010, 31 May 2011 and 4 July 2011 for 4 to 163 d and further exposed to a temperature elevation to 21°C in the laboratory. Day 0 corresponds to oysters not exposed to field conditions (i.e. control). The oysters used for OsHV-1 DNA detection were sampled at the end of the field exposure period, i.e. the onset of the temperature elevation to 21°C; nd: not detected

<table>
<thead>
<tr>
<th>Date of field exposure</th>
<th>Duration of exposure (d)</th>
<th>OsHV-1 DNA (log copies mg⁻¹)</th>
<th>No. of positive samples</th>
<th>Mortality</th>
</tr>
</thead>
<tbody>
<tr>
<td>22 November 2010</td>
<td>0</td>
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<td>0/2</td>
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</tr>
<tr>
<td></td>
<td>42</td>
<td>nd</td>
<td>0/2</td>
<td>No</td>
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<tr>
<td></td>
<td>73</td>
<td>nd</td>
<td>0/2</td>
<td>No</td>
</tr>
<tr>
<td></td>
<td>102</td>
<td>nd</td>
<td>0/2</td>
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</tr>
<tr>
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<td>135</td>
<td>nd</td>
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</tr>
<tr>
<td></td>
<td>149</td>
<td>nd</td>
<td>0/2</td>
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</tr>
<tr>
<td></td>
<td>162</td>
<td>1.7 ± 3.3</td>
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</tr>
<tr>
<td>31 May 2011</td>
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<td>6.1 ± 2.1</td>
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<td></td>
<td>17</td>
<td>6.9 ± 0.6</td>
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<td></td>
<td>20</td>
<td>8.5 ± 0.5</td>
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<td>4 July 2011</td>
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<td>nd</td>
<td>0/2</td>
<td>Yes</td>
</tr>
<tr>
<td></td>
<td>12</td>
<td>4.9 ± 0.5</td>
<td>2/2</td>
<td>Yes</td>
</tr>
</tbody>
</table>
Final survival of oysters increased from 30.9 to 61.4% in Aber Benoît and from 16.7 to 29.4% in the Bay of Brest in a manner consistent with oyster age (Fig. 3). The only exception was 9.9 mo old oysters, which exhibited relatively high mortalities at both sites.

In our study, age and size of oysters were positively correlated (whole body mass = 0.573 × age − 1.012; $r^2 = 0.953$). Therefore, the effect of oysters’ whole body mass on survival was similar to that of age (data not shown) and these 2 factors are confounded. However, there was no difference in size (shell length) between dead and live oysters measured after the onset of mortality in Aber Benoît on 15 June 2011, suggesting that disease mortality affected small and large animals similarly (Fig. 4; ANCOVA on shell length with age as a covariate and status as a fixed factor, status $p = 0.697$, age × status $p = 0.833$).

In our study, age and duration of field exposure were confounded factors. In a complementary experiment (see Supplement 2), we observed that survival clearly increased with age but slightly decreased with duration of field exposure. Therefore, it seemed that the effect of age was much more important than that of duration of field exposure.

Table 5. Expt C. Odds of disease mortality as a function of age of Pacific oysters *Crassostrea gigas* (varying from 4.3 to 11.2 mo old) in Aber Benoît and in the Bay of Brest for each time interval (0 to 136 d and 136 to 995 d). Custom hazard ratios were produced by means of polynomial (linear) exponentiated contrasts. The table also contains the standard error (SE) of the hazard ratio estimate and the confidence interval (CI) on the hazard ratio, the Wald $\chi^2$ statistic and the resulting p-value.

<table>
<thead>
<tr>
<th>Variable</th>
<th>Odds ratio</th>
<th>SE</th>
<th>95% CI</th>
<th>Wald $\chi^2$</th>
<th>p</th>
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</thead>
<tbody>
<tr>
<td>Age (linear)</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Aber Benoît</td>
<td></td>
<td></td>
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<td></td>
<td></td>
</tr>
<tr>
<td>0–136 d</td>
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<td>0.044</td>
<td>0.479–0.651</td>
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<td>&lt;0.001</td>
</tr>
<tr>
<td>136–995 d</td>
<td>1.072</td>
<td>0.207</td>
<td>0.735–1.565</td>
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<td>0.718</td>
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<td>Bay of Brest</td>
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<td></td>
<td></td>
</tr>
<tr>
<td>0–136 d</td>
<td>0.652</td>
<td>0.035</td>
<td>0.587–0.724</td>
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<td>136–995 d</td>
<td>1.362</td>
<td>0.188</td>
<td>1.039–1.785</td>
<td>5.0</td>
<td>0.025</td>
</tr>
</tbody>
</table>

Fig. 3. Expt C. Survival of Pacific oysters *Crassostrea gigas* exposed to field conditions in Aber Benoît or the Bay of Brest according to their age (in mo) at the onset of mortality, in relation to seawater temperature. Survival time was measured as days from the onset of mass mortalities in the Bay of Brest.
val increased with water renewal rate from 12.3 to 53.1% in exposed oysters and from 6.9 to 78.4% in challenged animals (Fig. 5A,B). The level of OsHV-1 DNA in exposed oysters at the onset of the experiment was $4.1 \times 10^6 \pm 3.6 \times 10^6$ copies mg$^{-1}$ wet tissue (n = 2 samples, data not shown), which is 2 orders of magnitude higher than the threshold value of $10^4$ DNA copies mg$^{-1}$ wet tissue at which OsHV-1 is considered to be involved in mortality (Oden et al. 2011, Schikorski et al. 2011a). After 2 and 4 d, the level of OsHV-1 DNA in exposed oysters increased, reaching $3.9 \times 10^7 \pm 3.9 \times 10^7$ copies mg$^{-1}$ wet tissue (n = 3 samples) and $2.4 \times 10^7 \pm 4.5 \times 10^7$ copies mg$^{-1}$ wet tissue (n = 4 samples) respectively. Survival of oysters varied as a function of status, biomass of infected oysters and time (Fig. 5C,D; Cox non-proportional hazard model, status $\times$ biomass $\times$ time $>[7 \text{ d}]$, Wald $\chi^2 = 17.5$, p < 0.001). In the first 7 d of the experiment, the parameter estimates for linear effects of biomass of infected oysters on survival of exposed animals was significant (Table 6). Based on this estimate, we calculated that an increase in the biomass of infected oysters to 108 g would increase the relative risk of mortality by 3.9 times in exposed animals. However, during this period, the biomass of infected oysters did not influence the risk of disease mortality in challenged oysters.

During the next period (7 to 19 d), this parameter estimate did not change in exposed oysters, but it increased markedly in challenged oysters (odds ratio = 10.2; Table 6). Therefore, the odds of mortality increased with the biomass of infected animals in both challenged and exposed oysters, but not at the same magnitude, nor at the same time.

Final survival decreased gradually with increasing biomass of infected animals from 91.2 to 74.6% in exposed oysters, and from 98.6 to 89.4% in challenged oysters (Fig. 5C,D). Mortalities were low in this experiment compared to the previous ones, likely reflecting that this experiment was conducted at the end of the infection period.

**DISCUSSION**

**Rearing history of oyster seed: natural spatfall vs. nursery**

Our results (Expt A) showed that 31/51 batches of oyster seed collected from the wild when seawater temperature was <14°C exhibited significant mortality under laboratory conditions at 21°C, compared to only 1/32 batches for animals sampled from a nursery. Therefore, the odds of mortality in wild oysters were 27 times higher than that of nursery animals. These results agree well with the fact that wild oyster seed is more exposed to disease risk in the field than those in nurseries, where they are likely to be protected from pathogens by means of prophylactic methods (Whittington et al. 2015).

We also observed that OsHV-1 µVar occurred at high prevalence in the wild oysters. They were presumably infected in spatfall collection sites when the seawater temperature was between 16 and 24°C (Pernet et al. 2012, 2014b, Dégremont 2013, Petton et al. 2013, Clegg et al. 2014). OsHV-1, as with other members of the *Herpesviridae* family, probably persists during the winter in its host after primary infection without inducing signs of disease or mortality (Arzul et al. 2002, Dundon et al. 2011, Dégremont et al. 2013).

Our results suggest that OsHV-1 is maintained in wild oysters in most French farming areas. As a result, it is most unlikely that OsHV-1 could be wiped out in sites where stocks of oysters are self-recruiting. However, in areas where oysters do not recruit, either due to low temperature or hydrodynamics, OsHV-1 could be cleared by ceasing restocking until the population dies off (Murray et al. 2012). In areas where OsHV-1 has not yet been detected, the movement of wild oysters from self-recruiting sites presents a high risk of disease transmission and mortality, compared to hatchery-
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Propagated seed and subsequent deployment in the field. Indeed, oysters collected from nurseries did not exhibit mortality at 21°C under laboratory conditions, and OsHV-1 DNA was not detected before or during the thermal challenge. Therefore, it appears that movement of wild oysters strongly increases the risk of disease transmission and mortality compared to that of nursery animals, which are generally safe before their deployment to the field until the temperature reaches ~15°C, a threshold temperature at which disease transmission occurs (see next section).

Fig. 5. Expt D. Survival of Pacific oysters *Crassostrea gigas* (A,C) exposed to field conditions in the Bay of Brest or (B,D) laboratory challenged (naive oysters placed in cohabitation with exposed animals) as a function of time. Exposed and naive oysters were placed in cohabitation at 3 different water renewal rates (48, 120 and 280 ml min⁻¹; A,B) or with 3 different biomasses of infected animals (50, 100 and 200 g; C,D)

Table 6. Expt D. Odds of disease mortality as a function of water renewal (48, 120 and 280 ml min⁻¹) and biomass of infected oysters (50, 100 and 200 g tank⁻¹) in exposed and challenged oysters for each time interval (0 to 7 and 7 to 19 d). Custom hazard ratios were produced by means of polynomial (linear) exponentiated contrasts. The table also contains the standard error (SE) of the hazard ratio estimate and the confidence interval (CI) on the hazard ratio, the Wald $\chi^2$ statistic and the resulting p-value

<table>
<thead>
<tr>
<th>Variable</th>
<th>Odds ratio</th>
<th>SE</th>
<th>95% CI</th>
<th>Wald $\chi^2$</th>
<th>p</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Water renewal (linear)</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Exposed 0−7 d</td>
<td>0.598</td>
<td>0.046</td>
<td>0.515−0.695</td>
<td>44.9</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>Challenged 0−7 d</td>
<td>0.532</td>
<td>0.103</td>
<td>0.364−0.777</td>
<td>10.6</td>
<td>0.001</td>
</tr>
<tr>
<td>Exposed 7−19 d</td>
<td>0.096</td>
<td>0.025</td>
<td>0.057−0.161</td>
<td>79.4</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>Challenged 7−19 d</td>
<td>0.208</td>
<td>0.063</td>
<td>0.115−0.375</td>
<td>27.2</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td><strong>Biomass (linear)</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Exposed 0−7 d</td>
<td>3.875</td>
<td>1.663</td>
<td>1.671−8.987</td>
<td>10.0</td>
<td>0.002</td>
</tr>
<tr>
<td>Challenged 0−7 d</td>
<td>1.28</td>
<td>0.168</td>
<td>0.989−1.656</td>
<td>3.5</td>
<td>0.061</td>
</tr>
<tr>
<td>Exposed 7−19 d</td>
<td>4.262</td>
<td>1.939</td>
<td>1.748−10.394</td>
<td>10.2</td>
<td>0.001</td>
</tr>
<tr>
<td>Challenged 7−19 d</td>
<td>10.275</td>
<td>4.92</td>
<td>4.020−26.264</td>
<td>23.7</td>
<td>&lt;0.001</td>
</tr>
</tbody>
</table>
Threshold temperature and onset of oyster mortality

The present study (Expts B and C) confirms that Pacific oyster mortality starts during spring when the seawater temperature reaches ~16°C, and coincides with detection of high quantities of OsHV-1 µVar DNA in oyster tissues (EFSA 2010, Segarra et al. 2010, Oden et al. 2011, Peeler et al. 2012, Pernet et al. 2012, 2014a, Dégremont et al. 2013, Petton et al. 2013, Clegg et al. 2014). It is, however, noteworthy that in Aber Benoît the onset of mortality occurred between 17 May and 15 June 2011, while average maximum daily seawater temperature was only 15.0°C. However during this time interval, seawater temperature had occasionally exceeded 16°C (the maximum seawater temperature recorded was 18.1°C on 15 June). Although the average daily seawater temperature of 16°C is a threshold above which disease transmission is optimal and mortalities occur, daily temperature variability likely enhances the risk of disease mortality, as previously shown in the withering syndrome in black abalone (Ben-Horin et al. 2013).

In contrast to Europe, OsHV-1 associated mortality of Pacific oysters in Australia occurs when seawater temperature is between 23 and 26°C (Jenkins et al. 2013, Paul-Pont et al. 2013, 2014). Therefore, it seems that the temperature threshold at which disease mortality occurs is not the same worldwide. The link between seawater temperature and the onset of mortality may be indirect, and it could involve other environmental, physiological or genetic factors that vary between Europe and Australia.

Seasonality of disease transmission: consequences for management of oyster movements

Healthy oysters deployed to the field between 22 November and 20 April when seawater temperature was <14.7°C exhibited no mortality in the laboratory at 21°C, and OsHV-1 DNA was not detected in their tissues, suggesting they were not infected by the virus (Expt A). Compared with previous studies which reported that no abnormal mortality occurred and OsHV-1 DNA was not detected in oyster tissues collected in the field during the winter (Oden et al. 2011, Pernet et al. 2012, Dégremont 2013), we further suggest that OsHV-1 is not transmitted to healthy oysters during this period.

In contrast, oysters left in the field until 3 May 2011 exhibited significant mortalities in the laboratory at 21°C and OsHV-1 DNA was detected in their tissues, when mortality had not yet been recorded in the field. Therefore, the temperature elevation under laboratory conditions revealed that OsHV-1 was transmitted to oysters between 20 April and 3 May, while daily average seawater temperature was between 14.7 and 15.8°C (the average seawater temperature during this period being 15.3 ± 0.3°C). Transmission of OsHV-1 under laboratory conditions from diseased to naïve oysters has been reported at temperature as low as 13.4°C and has led to significant mortalities in naïve animals at 14.4°C (Petton et al. 2013). Similarly in the Mediterranean Thau lagoon, OsHV-1 DNA was detected in sentinel oysters deployed at 33/104 locations when seawater temperature had just reached 16.0°C, indicating that disease transmission had occurred earlier, when seawater temperature was <16°C (Pernet et al. 2014b).

Oysters deployed in the field in spring (31 May) and summer (4 July) while seawater temperature was between 16 and 20°C showed significant mortalities in the laboratory at 21°C, concomitant with the detection of OsHV-1 DNA. This result corroborates that the optimal temperature range for disease transmission is between 16 and 22°C (Petton et al. 2013). Interestingly, survival of oysters in the laboratory at 21°C decreased with duration of field exposure in a manner similar to a dose–response relationship.

In France, oyster farming commonly involves the movement of large numbers of animals between rearing sites at successive steps in their production in order to optimize growth. These movements are likely to facilitate the spread of pathogens (EFSA 2010, Murray et al. 2012, Peeler et al. 2012, Murray 2013, Brenner et al. 2014). Our results show that the risk of infection of healthy oysters (in which OsHV-1 DNA was not detected and did not show any mortality after exposure at 21°C) exposed to field conditions was low, as long as daily average seawater temperature was <14.7°C, suggesting that further movements of these animals represents a low risk of disease transmission and mortality. However, when seawater temperature was between 14.7 and 15.8°C (~15.3°C), OsHV-1 was transmitted to oysters. Therefore, movements of healthy oysters during the spring (when no abnormal mortality has yet occurred) still presents a risk of infection and transmission of the virus. Finally, when seawater temperature is >16°C and mass mortalities have started, the risk of spreading OsHV-1 with oyster movements is high and increases with duration of exposure to field conditions where the virus is found.
Effects of age and size on disease susceptibility of oysters: consequences for seeding

In our study (Expt C), survival of oysters at 2 rearing sites increased markedly with age (and size) during disease outbreak. Similarly, other studies have shown that resistance of oysters to OsHV-1 increased with age and size (Dégremont 2013), likely reflecting maturation of the immune system (Green et al. 2014). These authors suggested that the host’s ontogeny influences the antiviral response in C. gigas. Alternatively, the higher cell replication rates of young animals could favour the replication of OsHV-1, as herpesviruses rely on the host’s cell replication machinery (Lyman & Enquist 2009).

The effect of age on the probability of mortality became progressively less important with time. This long-term effect may reflect that the cause of mortality had changed. For instance, we showed that the onset of mortality coincided with detection of OsHV-1 DNA by qPCR, although no pathogen analysis was conducted following detection. In support of this hypothesis, significant mortality occurred during the cold months, which is not compatible with OsHV-1 infection (EFSA 2010, Oden et al. 2011, Pernet et al. 2012, Petton et al. 2013).

The effects of oyster age and size on disease susceptibility were confounded in previous studies (Pernet et al. 2012, Dégremont 2013, Clegg et al. 2014, Paul-Pont et al. 2014). In our study as well, older oysters (which are less susceptible than younger ones) were larger. However, shell lengths of live and dead oysters after the outbreak of the disease were similar, suggesting that there was no relationship between disease susceptibility and oyster size. Similarly, Paul-Pont et al. (2014) reported that the age of oysters was a significant determinant of disease mortality rate, but size was not significant after allowing for variation in age. For example, these authors reported that 3 to 9 mo old oysters exhibited 80 to 100% mortality regardless of shell length, which varied from 2 to 96 mm (Paul-Pont et al. 2014).

Although survival of oysters varied as function of the interaction between time × site × age, survival of oysters increased as they aged both in Aber Benoît and in the Bay of Brest, as previously reported at Marennes-Oléron Bay (Dégremont 2013). In contrast, in the Mediterranean Thau lagoon, oysters remained highly susceptible to pathogen-related mortality pressure during their first 2 yr (Pernet et al. 2012). It is therefore possible that the age × site interaction affects mortality, as previously reported for the summer mortality syndrome (Dégremont et al. 2010b).

From a practical standpoint, our study has shown that the probability of mortality decreased with the age of oysters when facing the disease for the first time. Although the effect of age on odds of mortality became progressively less important with time, seeding older, healthy oysters led to higher overall survival. Therefore, it might be interesting to deploy oyster seed during the fall instead of the spring, so that animals are older and less susceptible when the disease occurs. This was shown for our healthy oysters, but may not be the case for asymptomatic carriers (as were 60% of our tested wild oyster batches; see Expt A).

Effect of water renewal and infectious biomass: perspectives for spatial planning

Under our controlled conditions, odds of disease mortality for oysters decreased markedly with water renewal and increased with the biomass of neighbouring infected animals, probably reflecting dilution and concentration effects respectively of viral particles in the seawater (Expt D). In the same way, survival of oysters cemented onto ropes, where circulation of water around each individual is enhanced and the density of susceptible hosts is lowered, was markedly higher (66%) compared to that of animals maintained in Australian baskets (13%), where the circulation of water around each individual and flushing rate was lower and the density of susceptible hosts was higher (Pernet et al. 2012).

Overall, these results corroborate that infectious disease dynamics are regulated by host population size and density, since larger and denser populations have increased contact between infectious and susceptible individuals (Anderson & May 1979). In terrestrial animal production, herd size and animal density are major risk factors for disease transmission (see e.g. Humblet et al. 2009 for review). This is also the case in finfish aquaculture (Krkošek 2010), where increased numbers of infectious fish lead to increased pathogen concentration in the surrounding environment, which in turn enhances the risk of pathogen transmission (Murray 2009). Similarly, a combined epidemiological–hydrodynamic model demonstrated that the number of infections in salmon increases with farm size (biomass) (Salama & Murray 2011).

The effects of water renewal and infectious biomass on disease transmission could explain the observed differences in oyster survival between and within farming sites. For instance, survival of oysters...
in the Thau lagoon, a semi-enclosed system in which the flushing rate is low (Fiandrino et al. 2003), is generally much lower than that observed along the Atlantic coast of France where animals are exposed to tidal regimes (Fleury & Bedier 2013). Such differences are not likely linked to temperature, as this parameter mainly triggers the timing of mortality rather than final survival (Petton et al. 2013). Additionally, survival of oysters within the bivalve farming area of the Thau lagoon (where the density of susceptible hosts is the highest) is much lower than outside it (Pernet et al. 2014b).

Consequently, spatial planning of oyster farming should take into account the hydrodynamic regime and biomass of the oysters, not only from a growth or a trophic perspective as is currently the case (Cranford et al. 2012), but also from an epidemiological viewpoint. Currently, the majority of oyster leases are concentrated in sheltered inshore areas where tidal currents are relatively low, which are favourable conditions for disease transmission and mortality. Due to space constraints, bivalve farms may move further offshore in higher tidal areas in order to allow the industry to expand. From an epidemiological perspective, offshore bivalve farming only makes sense if oysters are imported from areas where infection is not present.

**Thermal challenge of oysters: a complementary tool to OsHV-1 DNA detection by PCR**

In Expt A, we showed that detection of OsHV-1 DNA by real-time PCR in oysters sampled when seawater temperature was <14°C was not predictive of mortality risk. Although individual analyses of a greater number of oysters should be carried out to definitely rule out presence of OsHV-1, this result suggests that OsHV-1 DNA is hardly detectable in oysters when the virus is latent during the winter. However, the risk of mortality was highly correlated with detection of OsHV-1 DNA when oysters were exposed at 21°C. In the Thau lagoon, risk of oyster mortality was positively correlated with detection of OsHV-1 DNA in oysters while the seawater temperature was 16°C, but the odds ratio was only 1.4 (Pernet et al. 2014b). It is likely that the relationship between mortality risk and detection of OsHV-1 DNA is stronger when sampling time and the outbreak of disease are closer together, so that expression and detection of OsHV-1 can increase.

The lack of sensitivity of the OsHV-1 DNA PCR assay for detecting latent infection or asymptomatic carriers of the disease could become a limiting factor for developing proper disease management procedures. Indeed, improved diagnostic methods must be developed (EFSA 2010). Here, we provide evidence that thermal elevation of oysters to 21°C allows detection of asymptomatic carriers of OsHV-1. Our thermal challenge can be used for characterising the health status of oysters with regards to OsHV-1 complementary to PCR assays.

**CONCLUSIONS**

The present study provides evidence that the early rearing history, timing and duration of exposure to the disease in the field, age of oysters and water renewal and biomass of infected animals are important risk factors for OsHV-1 µVar transmission and related mortalities in *Crassostrea gigas*. Although some of these parameters were tested only under laboratory conditions, they are likely to have important implications for disease management in oyster farming systems. For instance, our thermal challenge can be used for characterising the health status of oysters with regards to OsHV-1 in conjunction with PCR assays. If the tested oysters show no mortality during the challenge and OsHV-1 DNA is not detected in oyster tissues, they can be qualified as ‘healthy’ and could be moved with a low risk of OsHV-1 transmission to other stocks. Additionally, increasing the age at deployment in the field will significantly increase oyster survival when OsHV-1 outbreaks occur (i.e. at seawater temperature >16°C). Finally, high hydrodynamics (flushing rate) and low biomass of infected animals in the environment will contribute to increased survival. If the tested oysters exhibit significant mortality associated with OsHV-1 during the challenge, they will potentially transmit the disease to neighbouring stocks when the seawater temperature reaches ~15°C, in a way similar to what we observed in Expt B. Infected animals will likely die when seawater temperature reaches 16°C in the field. Similar to healthy oysters, high hydrodynamic rates and low biomass of infected animals will mitigate mortality within the population. Although all tested parameters exerted a significant influence on the survival of oysters, their effects can be masked by the high infection pressure which currently occurs in most farming areas. For instance, 60% of natural seed batches tested in our study were infected by OsHV-1, suggesting that these oysters are likely to contribute to the global high infection pressure. Including the parameters explored in our study into
disease management scenarios may help to reduce the infection pressure of OsHV-1 in oyster farming systems.

Acknowledgements. The authors thank the Ifremer staff involved in oyster production at Argenton, the shellfish networks (Resco and Velyger) for providing temperature measurements, and Gaétan Daigle, Département de mathématique et statistique, Université Laval, Québec for advice with statistical analyses. This work was partly supported by Ifremer (project ‘Surmortalité’ 2010−11), the GIGASSAT project funded by ANR-AGROBiosphere N° ANR-12-AGRO-0001-01 and the French Ministry of Food, Agriculture and Fisheries (Convention DPMFA 2014 − Ifremer).

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Editorial responsibility: Marianne Holmer, Odense, Denmark

Submitted: July 28, 2014; Accepted: January 9, 2015
Proofs received from author(s): February 27, 2015