Salinity influences disease-induced mortality of the oyster *Crassostrea gigas* and infectivity of the ostreid herpesvirus 1 (OsHV-1)

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ABSTRACT: Mortality of young Pacific oysters *Crassostrea gigas* associated with the ostreid herpesvirus 1 (OsHV-1) is occurring worldwide. Here, we examined for the first time the effect of salinity on OsHV-1 transmission and disease-related mortality of *C. gigas*, as well as salinity-related effects on the pathogen itself. To obtain donors for OsHV-1 transmission, we transferred laboratory-raised oysters to an estuary during a disease outbreak and then back to the laboratory. Oysters that tested OsHV-1 positive were placed in seawater tanks (35‰, 21°C). Water from these tanks was used to infect naïve oysters in 2 experimental setups: (1) oysters acclimated or non-acclimated to a salinity of 10, 15, 25 and 35‰ and (2) oysters acclimated to a salinity of 25‰; the latter were exposed to OsHV-1 water diluted to a salinity of 10 or 25‰. The survival of oysters exposed to OsHV-1 water and acclimated to a salinity of 10‰ was >95%, compared to only 43 to 73% survival in oysters acclimated to higher salinities (Expt 1), reflecting differences in the levels of OsHV-1 DNA and viral gene expression (Expts 1 and 2). However, the survival of their non-acclimated counterparts was only 23% (Expt 2), and the levels of OsHV-1 DNA and the expression of 4 viral genes were low (Expt 1). Thus, OsHV-1 may not have been the ultimate cause of mortality in non-acclimated oysters weakened by a salinity shock. It appears that reducing disease risk by means of low salinity is unlikely in the field.

KEY WORDS: Bivalve · Herpesviridae · Infection · Disease transmission · Risk analysis

INTRODUCTION

Since 2008, massive mortality events of young Pacific oysters *Crassostrea gigas* have been reported in France (Miossec et al. 2009, EFSA AHAW 2010, Barbosa Solomieu et al. 2015, Pernet et al. 2016). These mortality events are associated with the infection of oysters with a newly described genotype (μVar) of ostreid herpesvirus 1 (OsHV-1) (Segarra et al. 2010). A causal relationship between OsHV-1 and oyster mortality has been established (Schikorski et al. 2011, EFSA AHAW 2015). This virus is now distributed along the European coastline from Portugal to Scandinavia, and closely related variants have been detected in Australia, New Zealand and Asia (Barbosa Solomieu et al. 2015, EFSA AHAW 2015, Pernet et al. 2016). OsHV-1 now poses a major challenge for Pacific oyster production around the world.

Oysters are grown in estuaries and bays, where they are often exposed to wide salinity fluctuations caused by evaporation, rainfall and inflow from rivers. Salinity may play a role in OsHV-1 transmission and disease development, as reported for other diseases of marine invertebrates (Haskin & Ford 1982, Ragone © The authors 2016. Open Access under Creative Commons by Attribution Licence. Use, distribution and reproduction are unrestricted. Authors and original publication must be credited. Publisher: Inter-Research · www.int-res.com
Most pathogens are susceptible to salinity because of their ionic permeability. For instance, decreasing salinity damages protozoan parasites because they are unable to regulate their cell volume (Ford & Haskin 1988, Burreson et al. 1994). Salinity also affects the occurrence and persistence of bacteria in the aquatic environment. For instance, the optimal salinity for the growth of pathogenic strains of *Vibrio splendidus* and *V. aegutianus* is approximately 20% (Vezzulli et al. 2015). To our knowledge, no information is available on the precise effects of salinity on eukaryotic marine viruses, but for viruses that infect bacteria, salinity influences viral adhesion to the hosts or genome injection by altering their envelope or capsid (Cordova et al. 2003, Kukkaro & Brussaard 2009, Mojica & Brussaard 2014).

The immune functions of bivalves are influenced by salinity with direct effects on the viability and activity of haemocytes (Matozzo & Marin 2011, Carregosa et al. 2014). Furthermore, the energy allocated for acclimation to salinity can impair defenses against pathogens (Gilles 1972, Shumway 1977, Shumway et al. 1977, Neufeld & Wright 1996). Thus, a change in salinity may increase host susceptibility to pathogens.

The aim of this study was to investigate the effect of salinity on disease transmission and the related mortality of *C. gigas*. Specifically, we examined the effect of salinity (1) on both the host and pathogens and (2) on the pathogens only. For both experiments, oysters were transferred to an estuary during a disease outbreak and then returned to the laboratory. Infected oysters were placed in tanks, and the surrounding seawater was used as a source of infection (SI). The SI was then (1) directed towards recipient oysters held at 4 salinities with or without prior acclimation or (2) brought to a salinity of 10 or 25‰ and directed towards recipient oysters maintained at 25‰. Overall, this method, which has been used in many previous studies (Petton et al. 2013, 2015a,b, Tamayo et al. 2014, Lassudrie et al. 2015, Lemire et al. 2015, Pernet et al. 2015), reproduces the natural infection process and takes into account the fact that OsHV-1 triggers oyster mortality, although other microorganisms can also play a role (Saulnier et al. 2010, Lemire et al. 2015, Petton et al. 2015b).

**MATERIALS AND METHODS**

**Animals**

The 4 mo old oysters used in Expts 1 and 2 were taken from 2 cohorts, produced in 2014 and 2015, respectively, according to Petton et al. (2015a). Adults originating from Fouras (France, 46°00'43.2'' N, 1°07'02.9” W) were moved to the Ifremer marine station in Argenton (48°48’24.49” N, 3°00’22.84” W) for conditioning in January 2014 and March 2015 for Expts 1 and 2, respectively. These oysters were maintained at 17°C in 500 l flow-through tanks. Fertilization was achieved on 11 March 2014 (Expt 1) and on 28 April 2015 (Expt 2) by stripping the gonads. The embryos developed in 150 l tanks at 21°C for 48 h, and D-larvae were transferred to flow-through rearing systems at 25°C. After 15 d, competent larvae were collected and allowed to settle in downwellers. The entire rearing cycle was conducted in UV-sterilized, 1 μm filtered seawater enriched with living phytoplankton (Petton et al. 2015a). At the onset of both experiments (17 July 2014 and 7 August 2015), the total body mass of oysters was similar at 0.8 ± 0.2 g (mean ± SD).

During the experiments, oysters were fed with a mixture of *Chaetoceros muelleri* (CCAP 1010/3) and *Tisochrysis lutea* (CCAP 927/14) (1:1 in dry weight). Food concentration was ca. 1300 μm³ μl⁻¹ of microalgae at the outlet of the rearing tanks. Seawater was sampled daily at the inlet and outlet of each experimental tank to determine phytoplankton consumption. Cell concentrations were measured using an electronic particle counter (Multisizer 3) equipped with a 100 μm aperture tube. Salinity, temperature, pH and oxygen were controlled daily with the WTW probes cond340, xi3101, pH3310 and FDO 925, respectively. Each tank was lightly bubbled to maintain oxygen levels between 85 and 100% of saturation.

**Method of infection**

The method of infection was a modified procedure from Petton et al. (2015a). Oysters were transferred on 22 July 2014 (Expt 1) and 6 August 2015 (Expt 2) to a farming area of the Bay of Brest (48°20’06.19” N, 4°19'06.37'' W) where oyster mortality was occurring. The salinity was 34‰. After a week of exposure to field conditions, few dead oysters were observed, and the remaining live individuals were moved back to the Ifremer facilities in Argenton. These oysters tested positive for the OsHV-1 μVar strain, and the
level of virus DNA was $1.9 \times 10^9 \pm 1.6 \times 10^9$ copies mg$^{-1}$ fresh weight (mean ± SD; n = 3 pools of 15 oysters), they were therefore considered useful as OsHV-1 donors and placed in 45 l flow-through tanks at 35‰ and 21°C (Petton et al. 2013). The seawater surrounding the donors was used as the SI.

For Expt 1, the SI was connected by flexible tubes fitted inside a peristaltic pump to 45 l tanks (n = 12), which contained the recipient oysters at 4 salinities. The other 12 tanks were not connected to the SI and were used as controls. For each tank, the water flow from the SI was 11% of the total water flow (12 l h$^{-1}$). The water input from the SI was arrested on Day 6, when the recipients exhibited mortality. For Expt 2, the SI was connected to 45 l tanks (n = 6), which contained water at either 10 ± 0.7‰ (low-salinity tanks, n = 3) or 25.1 ± 0.4‰ (control tanks, n = 3). These salinities were obtained by mixing seawater from the SI (35‰) with fresh water. These tanks were then connected to 45 l tanks (n = 6) containing recipient oysters maintained at 25.4 ± 0.2‰. In each tank, the water flow from the SI was maintained at 11% of the total water flow fixed at 6 l h$^{-1}$ for 4 d.

**Experimental design**

We first investigated the effect of 4 salinities (mean ± SD: 10.2 ± 0.4, 15 ± 0.8, 24.9 ± 0.4, 35.4 ± 0.2‰) encountered in the field (Fig. 1) on disease transmission and the subsequent mortality of Pacific oysters (Expt 1). These salinities were obtained by mixing 1 μm filtered and UV sterilized seawater (35‰) with fresh water from the tap (chlorine level <0.3 mg l$^{-1}$). Between 17 and 28 July 2014, oysters were acclimated to each salinity in 45 l tanks (Fig. 2, n = 6 tanks, 310 oysters tank$^{-1}$). Filtration rate, i.e. the volume of microalgae consumed per gram of oyster per day, was measured daily in each tank using the following formula:  

$$\text{Filtration rate (μm}^3 \, \text{g}^{-1} \, \text{d}^{-1}) = \frac{([\text{Cell}_{\text{inlet}}] - [\text{Cell}_{\text{outlet}}] \times \text{Waterflow})}{\text{Biomass of oysters}}$$

where the variables were the concentrations of microalgae at the inlet and outlet of the tank ([Cell$_{\text{inlet}}$] and [Cell$_{\text{outlet}}$] in μm$^3$ of algae per μl of seawater); the water flow in the rearing tank, which was maintained at 12 l h$^{-1}$; and the biomass of oysters in each tank. On 28 July 2014 (Day 0), the biomass of acclimated

Oysters was standardized at ca. 160 g in each tank to remedy differences in the growth rates among the salinity treatments. In the same tanks, 160 g of non-acclimated oysters were added. Half of the tanks were connected for 6 d to the SI, and the other half were used as controls. Survival of recipient oysters was monitored daily for 17 d, and dead oysters were removed from tanks.

We then investigated the effect of salinity on the infectivity of OsHV-1 (Expt 2). Between 7 and 13 August 2015, oysters were acclimated at 25‰ in 45 l tanks (Fig. 2, n = 6 tanks, 180 oysters tank⁻¹). On 13 August 2015 (Day 0), the biomass of acclimated oysters was standardized at ca. 150 g of total oyster biomass in each tank. The tanks were connected to the SI. Survival of recipient oysters was monitored daily for 17 d, and dead oysters were removed from tanks.

Alive recipient oysters (15 for Expt 1 and 10 for Expt 2) were sampled in each tank on Days 0, 2 and 4 before the onset of mortality (Petton et al. 2013, 2015a). Whole tissues were removed from the shells, pooled together, flash frozen in liquid nitrogen and stored at −80°C. The oyster tissues were then ground in liquid nitrogen with a MM400 homogenizer (Retsch), and the resulting oyster powder was subsampled (30 mg of fresh weight) for quantification of OsHV-1 DNA and viral gene expression.

OsHV-1 DNA quantification

These analyses were conducted by the Laboratoire Départemental Vétérinaire de l’Hérault (Montpellier, France) using oyster powder homogenized in sterile artificial seawater (Pepin et al. 2008). Total DNA was then extracted with a QIAamp tissue mini kit (Qiagen) according to the manufacturer’s protocol. The extract was stored at −20°C before detection and quantification according to a real-time PCR protocol based on SYBR® Green chemistry (Pepin et al. 2008) with specific primers validated by Webb et al. (2007).

The results were expressed as the number of OsHV-1 DNA copies per mg of wet tissue.

OsHV-1 gene expression

Total RNA was extracted with EXTRACT-ALL® (Eurobio) according to the manufacturer’s recommendation, and 20 µg of extracted RNA were then removed from the genomic DNA using a DNase Max™ Kit (MO Bio). The RNA quantity and quality were determined with a NanoDrop 2000 spectrophotometer (Ambion®). A No-RT was then performed to confirm the absence of genomic DNA. First-strand cDNA synthesis was performed using the SuperScript III First-Strand Synthesis System (Invitrogen) on 500 ng of treated RNA.

The OsHV-1 genome has 207 439 bp and contains 124 unique genes (Davison et al. 2005). In this study, open reading frames 72, 75, 87 and 117 were selected from Segarra et al. (2014). These genes encode for a membrane protein, a dUTPase enzyme, an apoptosis inhibitor and a ring finger protein (Davison et al. 2005, Segarra et al. 2014). Gene expression was expressed as a delta threshold cycle number (Ct) and calculated using the following formula: Ct viral gene − Ct house-keeping gene (elongation factor).

Real-time (RT) quantitative PCRs were performed with an Mx3005P Thermocycler sequence detector (Agilent). The forward and reverse primers used for this study were described by Segarra et al. (2014). Amplification reactions were done in duplicate in a total volume of 20 µl. Each well contained 5 µl of cDNA solution (1/30, diluted in nuclease-free water), 10 µl of Brilliant® SYBR® Green III PCR mix (Agilent), 2 µl of forward and reverse primers (3 µM) and 1 µl of distilled water. The thermal cycling conditions for the PCRs were 3 min at 95°C followed by 40 amplification cycles at 95°C for 5 s and 60°C for 20 s. To ensure the specificity of the primers used, melting curves were plotted (55–95°C). For each qPCR,
negative controls were included to rule out DNA contamination.

**Statistical analyses**

**ANOVA**

The differences in oyster total body mass at the end of the acclimation period among the salinity treatments were analysed by ANOVA (Expt 1). Mixed-design ANOVAs were performed to assess differences in (1) the filtration rate depending on salinity (4 levels, main plots) and acclimation time (8 levels, subplots); (2) the levels of OsHV-1 DNA and viral gene expression in oysters, depending on (1) salinity and infection (4 × 2 levels, main plot), acclimation (2 levels, subplots), time (for DNA only, 2 levels, subsubplots) (Expt 1) and (2) salinity (2 levels, main plots) and time (2 levels, subplots) (Expt 2). Differences in viral gene expression among the salinity treatments were analysed by ANOVA (Expt 2).

The unit of replication was the tank in which the salinity and infection treatments were applied. All mutual interactions among the factors were tested, and Tukey’s HSD was used as a post hoc test. The normality of residuals and homogeneity of variances were graphically checked, and the data were log \((x + 1)\) transformed where necessary. ANOVAs were carried out using R (www.R-project.org/).

**Survival analysis**

Nonparametric estimates of the survivor function were computed according to Kaplan & Meier (1958). Survival time was measured as days from the onset of the experiment (Day 0) when oysters were exposed to the SI. The data were read as the number of dead animals within each tank on each day. Survival curves of exposed oysters were plotted and compared among the salinity treatments.

In Expt 1, the survival time curves of oysters connected to the SI were compared using the Cox regression model (Cox 1972) after adjustment for the effect of salinity and acclimation. The survival of control oysters (not exposed to the SI) was not included in the statistical models because it was always >95%. 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: \(t \leq 6\) d or \(t > 6\) d. Custom hazard ratios were produced by means of contrasts. Cox regression was conducted using SAS 9.4 (SAS Institute).

**RESULTS**

**Salinity modulates host–pathogen interaction (Expt 1)**

Prior to the experiment, the filtration rates of oysters at 10 and 15‰ were initially lower than those of oysters at 25–35‰. The rates then increased during the first 2 to 6 d of acclimation until reaching the filtration rates of oysters at 25–35‰ (Fig. 3A, Table S1 in the Supplement at www.int-res.com/articles/suppl/q008p543_supp.pdf). At the end of the acclimation period, the total body mass of oysters at different salinity levels was ranked as follows: 10 < 15 < 25 = 35‰ (Fig. 3B, Table S2 in the Supplement). At 10‰, the oysters did not grow (\(p = 0.103\)).

Survival of control oysters was 99.1 ± 1.9% (mean ± SD) irrespective of salinity and acclimation treatments. Consequently, only oysters exposed to the SI were considered in this section.

At the end of the experiment, the survival of oysters exposed to the SI and acclimated at 10‰ was higher (95.8%) than that of oysters held at 15, 25 and 35‰,
where survival was 73.2, 43.2, and 61.9%, respectively (Fig. 4A). In contrast, when oysters were exposed to the SI without prior acclimation to the salinity, their survival at 10‰ was markedly lower (22.8%) than at 15, 25, and 35‰, where survival was 63.5, 42.2, and 48.5%, respectively (Fig. 4B).

During the first 6 d, the oyster mortality hazard was ranked as 10 < 15 < 35 < 25‰ or 10 < 15 < 25 = 35‰ for acclimated and non-acclimated oysters, respectively (Table S3 in the Supplement). Between 7 and 17 d, the mortality hazard in acclimated oysters remained fairly similar (10 < 35 < 15 < 25‰) to that observed between 0 and 6 d. Overall, low salinity decreased the risk of mortality of acclimated oysters exposed to the SI. This was not the case in non-acclimated oysters, where the mortality hazard at 10‰ was 3 to 4 times higher than that at 15–35‰ between 7 and 17 d (Table S3).

On Day 0, OsHV-1 DNA was not detected in recipient oysters. Salinity influenced the levels of OsHV-1 DNA in exposed oysters but not in controls on Days 2 and 4 (Fig. 5A, Table S4 in the Supplement). In fact, at 10‰, the level of OsHV-1 DNA in exposed oysters was as low as in the controls (4.7 \times 10^3 copies mg\(^{-1}\)), but it reached more than 10^6 copies mg\(^{-1}\) at higher salinities. Viral gene expression in exposed oysters was ranked as 10 < 15 = 25 = 35‰, and at 10‰ it was as low as in the controls (Fig. 5B, Table S5 in the Supplement).

Low salinity reduces OsHV-1 infectivity (Expt 2)

When the SI was treated for 7 h at 10‰ before coming into contact with oysters at 25‰, survival of the oysters remained 100% (Fig. 6A). In contrast, when the SI was treated at 25‰ (control), survival decreased to 44.8% after 14 d. The level of OsHV-1 DNA in oysters exposed to the SI treated at 10‰ was markedly reduced compared to that of controls (Fig. 6B, Table S6 in the Supplement). The levels of viral gene expression in oysters after the SI was treated at 10‰ were lower than that expressed after exposure to 25‰ (Fig. 6C, Table S7 in the Supplement).

**DISCUSSION**

The survival of oysters exposed to the SI and acclimated at 10‰ was always >95%, compared to only 43 to 73% at higher salinities. Concomitantly, the levels of OsHV-1 DNA and the expression of 4 viral genes in oysters were lower at 10‰ than at other salinities. Therefore, the high survival at 10‰ coincided with low levels of OsHV-1 DNA and viral gene expression. Although the effect of salinity on virus–bivalve interactions has never been investigated, it is known to influence other host–pathogen interactions in the marine environment. Indeed, low salinity reduces the disease risk and prevalence of protozoan parasites in oysters and clams (Andrews & Wood 1967, Haskin & Ford 1982, Ragone Calvo et al. 1998, Brothers et al. 2000, Soudant et al. 2013). We further investigated whether the increased survival of oysters acclimated at 10‰ exposed to the SI reflects changes in pathogen infectivity or in host physiology or both.

Low salinity (10‰) clearly reduced OsHV-1 infectivity because (1) the survival of oysters held at 25‰...
and further exposed to the SI previously treated at 10‰ was 100%, compared to only 45% in oysters exposed to the SI treated at 25‰, and (2) the levels of OsHV-1 DNA and the viral gene expression in oysters were much lower when the SI was treated at 10‰. Studies show that salinity influences the infectivity of bacteriophages by altering their structures and changing the adsorption rate to host cells (Mojica & Brussaard 2014). It is likely that OsHV-1, as an enveloped virus (Renault et al. 2004, Davison et al. 2005), is particularly susceptible to changes in envi-

Fig. 5. (A) Quantification of OsHV-1 DNA, expressed as DNA copy numbers per mg of fresh weight and (B) virus gene expression as a delta threshold cycle number (Ct) of OsHV-1 open reading frames (ORFs) relative to Pacific oyster *Crassostrea gigas* elongation factor in live oysters as a function of salinity, exposure to the source of infection and acclimation. Data are means ± SE (n = 3 tanks). For OsHV-1 DNA, data were log (x+1) transformed and are the means of Days 2 and 4. The viral gene expression analyses were conducted on Day 2 only. Different letters indicate significant differences.

Fig. 6. (A) Survival of Pacific oysters *Crassostrea gigas*, (B) quantification of OsHV-1 DNA, expressed as DNA copy number per mg of fresh weight and (C) virus gene expression as a delta threshold cycle number (Ct) of OsHV-1 open reading frames (ORFs) relative to *C. gigas* elongation factor in live recipient oysters maintained at 25‰ and exposed to the source of infection previously treated at 10 and 25‰. Survival time was measured as days from the onset of exposure to the source of infection. Data are means ± SE (n = 3 tanks). For OsHV-1 DNA, the data were log (x+1) transformed and are the means of Days 2 and 4. The gene expression analyses were conducted on Day 2 only. Different letters indicate significant differences.
ronmental parameters such as salinity. This type of virus is generally more susceptible to environmental changes than non-enveloped viruses (Lucas 2001, Kukkaro & Bamford 2009).

Prior acclimation to salinity plays a major role in the host response to a pathogen. The survival of oysters exposed to the SI and acclimated at 10% was always >95%, whereas the survival of their non-acclimated counterparts held in the same tanks was only 23%. Nevertheless, the levels of OsHV-1 DNA and viral gene expression in these non-acclimated oysters were similar to those of controls and acclimated animals that showed no significant mortality. Therefore, OsHV-1 may not be the cause of mortality in these oysters. A more likely hypothesis is that another microorganism killed these animals simultaneously weakened by a salinity shock. In our study, donors were infected in the field where the OsHV-1-induced mortality occurred, and they may have introduced a diversity of Vibrio species or other pathogens to the recipients (Lemire et al. 2015, Petton et al. 2015b). In addition, acute exposure to low salinity may have altered the defenses of oysters either directly by impairing their immunity (Matteo & Marin 2011, Carregosa et al. 2014, Knowles et al. 2014) or indirectly by causing the oysters to allocate energy to regulate intracellular osmotic concentrations (Gilles 1972, Shumway 1977, Shumway et al. 1977, Neufeld & Wright 1996), thus compromising the defense response.

Oyster mortality began after only 2 to 3 d of exposure to the SI except for the non-acclimated animals at 10 and 15%, in which it occurred after 6 d. This delay may reflect salinity acclimation time. Indeed, bivalves exposed to an acute change in salinity first close their shells and reduce feeding (Pierce & Berggreen 1972). This behaviour was observed during acclimation prior to the experiment. Considering that suspension feeding is the portal of entry for pathogens in bivalves (Ben-Horin et al. 2015), the reduction in the oyster feeding rate at low salinity may have decreased the pathogen exposure, leading to delayed mortality.

At the onset of infection, oysters acclimated to different salinities showed major differences in growth, but their levels of OsHV-1 DNA and viral gene expression after 2 d were similar. Therefore, it seems that the growth rate of oysters did not influence OsHV-1 replication. This result may appear somewhat paradoxical with the idea that the host cell machinery modulates the replication of herpes viruses (Lyman & Enquist 2009). More fundamental information on the physiology of the host with respect to viral infection is needed. For instance, the impact of oyster growth conditions on their susceptibility to the virus and on the rate of virus proliferation has never been studied.

Differences in oyster survival at 15, 25 and 35% with similar levels of OsHV-1 DNA and viral gene expression may reflect differences in bacterial communities. Indeed, several non-pathogenic bacteria contribute to the mortality of oysters (Lemire et al. 2015), and salinity is a major factor shaping the bacterial community in marine ecosystems (Herlemann et al. 2011, Dupont et al. 2014).

Estuaries and coasts, the natural habitats of oysters, are generally subject to wide fluctuations in salinity. However, salinities lower than 15% occur for only a few hours or days under natural field conditions in France (Fig. 1). Therefore, it appears that oysters are rarely acclimated to low salinity in the field, and reducing disease risk by means of low salinity is unlikely.

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