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# **Research Article**

Cite this article: O' Reilly AJ, Laide C, Maloy A, Hutton S, Bookelaar B, O' Sullivan K, Lynch S A, Culloty SC (2018). The role of the mussel *Mytilus* spp. in the transmission of ostreid herpesvirus-1 microVar. *Parasitology* **145**, 1095–1104. https://doi.org/10.1017/S0031182017002244

Received: 27 April 2017 Revised: 3 November 2017 Accepted: 14 November 2017 First published online: 21 December 2017

#### Key words:

Cohabitation; Crassostrea gigas; Herpesvirus; interspecies transmission; Mytilus spp.; temperature

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# The role of the mussel *Mytilus* spp. in the transmission of ostreid herpesvirus-1 microVar

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#### **Abstract**

The Pacific oyster Crassostrea gigas contributes significantly to global aquaculture; however, C. gigas culture has been affected by ostreid herpesvirus-1 (OsHV-1) and variants. The dynamics of how the virus maintains itself at culture sites is unclear and the role of carriers, reservoirs or hosts is unknown. Both wild and cultured mussels Mytilus spp. (Mytilus edulis, Mytilus galloprovincialis and hybrids) are commonly found at C. gigas culture sites. The objective of this study was to investigate if Mytilus spp. can harbour the virus and if viral transmission can occur between mussels and oysters. Mytilus spp. living at oyster trestles, 400-500 m higher up the shore from the trestles and up to 26 km at non-culture sites were screened for OsHV-1 and variants by all the World Organization for Animal Health (OIE) recommended diagnostic methods including polymerase chain reaction (PCR), quantitative PCR (qPCR), histology, in situ hybridization and confirmation using direct sequencing. The particular primers that target OsHV-1 and variants, including OsHV-1 microVar (µVar), were used in the PCR and qPCR. OsHV-1  $\mu$ Var was detected in wild Mytilus spp. at C. gigas culture sites and more significantly the virus was detected in mussels at non-culture sites. Cohabitation of exposed wild mussels and naïve C. gigas resulted in viral transmission after 14 days, under an elevated temperature regime. These results indicate that mussels can harbour OsHV-1 µVar; however, the impact of OsHV-1  $\mu$ Var on *Mytilus* spp. requires further investigation.

# Introduction

The Pacific oyster Crassostrea gigas and mussels Mytilus spp. are commercially important shellfish species, which have contributed to the substantial growth of the aquaculture industry in Ireland. Both wild and cultured mussels, Mytilus edulis, Mytilus galloprovincialis and hybrids of both species, are distributed along the Irish coastline (Lynch et al. 2014) and are often located in close proximity to cultured C. gigas stocks. Ostreid herpesvirus-1 (OsHV-1) and variants such as OsHV-1 var. and OsHV-1 microVar (µVar) belong to the family Malacoherpesviridae, the genus Ostrea virus containing the species OsHV-1 (Davison et al. 2009). In recent years, OsHV-1  $\mu$ Var have been associated with C. gigas mortalities in several countries worldwide, including France (Segarra et al. 2010), Australia (Jenkins et al. 2013), Italy (Dundon et al. 2011), Spain (Roque et al. 2012) and Ireland (Peeler et al. 2012), with an additional Irish genotype also being identified (Lynch et al. 2012). In Ireland, C. gigas mortalities were first recorded in both adult and spat oysters in 1993 and were associated with OsHV-1 (Cotter et al. 2010). OsHV-1 µVar was first identified in Ireland in 2008 following mortality events in three separate bays and by 2012 in a total of 29 bays around the Irish coastline (Morrissey et al. 2015). The herpesvirus combined with suboptimal environmental factors forms the basis of high economic risk for Pacific oyster culture (Paredes et al. 2013).

Herpes-like virus infection has been observed in bivalve species such as the flat oyster *Tiostrea chilensis* in New Zealand (Hine *et al.* 1998); *Ostrea angasi* in Australia (Hine and Thorne, 1997); European flat oyster, *Ostrea edulis* in France (Comps and Cochennec, 1993; Renault *et al.* 2000); Manila clam, *Ruditapes philippinarum* (Renault, 1998; Renault and Arzul, 2001; Renault *et al.* 2001); scallop (Arzul *et al.* 2001) and abalone *Haliotis diversicolor supertexta* (Chang *et al.* 2005; Tan *et al.* 2008). OsHV-1 μVar DNA was detected in wild Mediterranean mussels *M. galloprovinciallis* in California and in cultured *M. galloprovinciallis* in Italy (Burge *et al.* 2011; Domeneghetti *et al.* 2014). More recently OsHV-1 μVar DNA was detected in the Sydney rock oyster *Saccostrea glomerata*, Sydney cockle *Anadara trapezia*, blue mussels *Mytilus spp.*, hairy mussel *Trichomya hirsuta*, whelks *Batillaria australis* and barnacles *Balanus spp.* in Australia (Evans *et al.* 2017). Filter feeding bivalves such as mussels, oysters and clams may typically harbour various pathogenic organisms such as Herpes and enteric viruses (Venier *et al.* 2011), as bivalves process large volumes of water during feeding thus maximizing their exposure (Witte *et al.* 2014).

Burge et al. (2011) quantified OsHV-1 viral loads in C. gigas and wild M. galloprovincialis in California and viral copy numbers per ng of genomic DNA of OsHV-quantitative

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polymerase chain reaction (qPCR) in positive individuals observed was much lower in M. galloprovinicialis (0·0073  $\pm$  0·006; 0·0005 to 0·025) compared with C. gigas (20·31  $\pm$  12·40; 0·005 to 114·4). Domeneghetti et al. (2014) reported that OsHV-1  $\mu$ Var in C. gigas and M. galloprovinciallis, co-cultured in an Italian lagoon with seawater temperatures of 17·6 °C, had lower levels of viral DNA in the asymptomatic mussels compared with the oyster spat (up to 8·4  $10^4$ /ng DNA) sampled before an acute mortality event. OsHV-1  $\mu$ Var was detected in asymptomatic mussels cohabiting with the virus-positive oysters and these M. galloprovincialis had a mean load of 3·9 virus copies/10 ng of DNA (Domeneghetti et al. 2014). Although OsHV-1  $\mu$ Var DNA was detected in M. galloprovinicialis in both of those studies, transmission of the virus from mussels to oysters and vice versa was not investigated.

Elucidating the role of mussels in viral maintenance and transmission, i.e. their ability to act as carriers, reservoirs or incidental hosts, is an important component in understanding viral dynamics and in developing risk management measures. A host that becomes infected, but is not required for the maintenance of the population of a pathogen, can be termed (a) an incidental host (Ashford, 2003). Haydon et al. (2002) defined a (b) reservoir host as 'one or more epidemiologically connected populations or environments in which the pathogen can be permanently maintained and from which infection is transmitted to the defined target population'. A (c) 'carrier' is an individual with no overt disease, which harbours infectious organisms (Brachman, 1996). A (d) viral reservoir has been defined 'as a cell type in which a replication-competent form of the virus accumulates and persists with more stable properties than the main pool of actively replicating virus' (Blankson et al. 2002). Reservoirs of infection can be comprised of one or more interacting species (Haydon et al. 2002 cited in Viana et al. 2014).

The direct relationship between viral proliferation, transmission and environmental stressors has been demonstrated particularly temperature influence and increasing temperatures associated with oyster mortality, in field studies (Garcia *et al.* 2011) and under experimental conditions (Sauvage *et al.* 2009; Pernet *et al.* 2012, 2015; Petton *et al.* 2013; Martenot *et al.* 2015). Oyster mortality events have followed increases in seawater temperature exceeding 19 °C (Pernet *et al.* 2010). Pernet *et al.* (2015) observed that OsHV-1  $\mu$ Var was persistent in oysters held at a low temperature of 13 °C, with no associated mortalities, but the virus was reactivated during thermal elevation to 21 °C, thus highlighting that mortality in OsHV-1  $\mu$ Var infected seed oysters was only delayed if those oysters were moved to low-temperature location because once temperatures increased the virus was activated and oyster mortalities occurred.

The study of viruses and the role of reservoirs is fundamental for a better understanding of the intrinsic relationship between the virus and host (Appolinário et al. 2015). It is important to better manage the risks associated with this virus and knowing which additional carriers, hosts or reservoirs are involved in the maintenance of the virus is imperative for the implementation of future management strategies of disease control. Other studies have highlighted the importance of this including studies on the protistan pathogen Bonamia ostreae of the European flat oyster O. edulis, which highlighted the detection of this pathogen in non-typical hosts, such as C. gigas (Lynch et al. 2010) and benthic macroinvertebrates and zooplankton (Lynch et al. 2007). Implementing precautionary measures when moving shellfish consignments and non-typical hosts from pathogen infected to uninfected areas will benefit the control and management of such pathogens, including OsHV-1 µVar. The objectives of this study were (a) to determine if OsHV-1 µVar was present in

Irish wild *Mytilus* spp. living in close proximity to oyster trestles, approximately 400–500 m from the oyster culture sites and up to 26 km from the oyster farms at non-culture sites and (b) investigate transmission dynamics and viral detection in mussels between naturally exposed mussels and naïve oysters under an elevated temperature regime in the laboratory. Findings from this study will contribute to a more comprehensive understanding of the underlying relationship between viral proliferation among cultivated oysters and mussel species and provide an insight into the interspecific species interactions and transmission that occurs in a marine environment.

### Materials and methods

Study sites and field sampling

As the sites used in the study contain a mixture of *M. edulis*, *M. galloprovincialis* and hybrids mussels (with the exception of Carlingford Lough where *M. edulis* is exclusively found), all the following descriptions refer to *Mytilus* species (spp.). Wild mussels from both oyster culture and non-culture (control) sites were screened.

# Field survey 1

A preliminary field survey, to determine if OsHV-1  $\mu$ Var could be detected in mussels from hybrid zones (M. edulis, M. galloprovincialis and hybrids of both parent species) in Ireland took place in 2014 with (a) a single mussel sample (n = 30) being collected in late June from Goat Island, approximately 5 km west of Ardmore, County Waterford (51°57′04"N, 7°43′23"W) and approximately 26 km from a OsHV-1 μVar endemic C. gigas culture site (Table 1), (b) a single mussel sample (n = 250) was collected in July from Garrettstown, Kinsale, Co. Cork (51° 38'16.93"N, 8°34'21.41"W), which is approximately 22 km from a OsHV-1  $\mu$ Var endemic C. gigas culture site from which (n =30) mussels were initially screened. An additional Garrettstown mussel sample (n = 30) was screened after the mussels collected in July were held in the laboratory for 3 weeks and held at 13 °C and (c) several mussel samples were collected in July on the south east coast of Ireland at Ballymacoda Bay, Youghal Co. Cork (51°53′24"N, 7°55′57"W). At that site, wild mussels were sampled at a rocky outcrop on the mid to lower intertidal approximately 400 m from a C. gigas OsHV-1 µVar designated culture site, a site where significant oyster mortalities occurred in 2009 and 2010 (www.marine.ie/Home/site-area/news-events/ news/update-oyster-mortalities). An initial sample (n = 42) of adult-sized mussels Mytilus spp. was collected in mid-July and a second random sample (n = 167) of Mytilus spp. was collected in late-July (Table 1).

## Field survey 2

Further sampling and screening of mussels took place as part of a second larger field study between 2013 and 2015 at two *C. gigas* culture sites: (d) Carlingford Lough (54·0733° N, 6·1994° W), on the east coast of Ireland in the Irish Sea and (e) Dungarvan Harbour (52°03′ N, 007°35′ W), on the south east coast in the Celtic Sea. OsHV-1 µVar is endemic at both of these sites since 2009 (www.marine.ie/Home/site-area/news-events/news/update-oyster-mortalities). Dungarvan is a known hybrid zone, while only *M. edulis* is found at Carlingford Lough. At both culture sites, mussels were collected from the *C. gigas* PVC culture bags at the oyster trestles on the mid to lower intertidal and approximately 500 m from the trestles at the high shore. At Carlingford Lough, sampling took place on nine occasions, every 2 weeks from July to October in 2013 and on eight occasions, every 2 weeks from April to

Table 1. Summary of Mytilus spp. samples collected at five Irish sites in field surveys 1 and 2 in 2013-2015

Sample date	Sample site	No. of individuals and shore height	Sample type	OsHV-1 $\mu$ Var prevalence (%)	Mean max–mean min. (mean daily) temperature (°C) <sup>a</sup>
Field survey 1					
29/06/2014	Ardmore	n = 30	Gill	7 (2/30)	16·9–9·9 (13·4)
01/07/2014	Garretstown	n = 250	Gill	25 (15/60)	19·8–10·1 (15·0)
20/07/2014	Ballymacoda initial sample	n = 42	Gill, haemolymph, shell cavity fluid	100 (12/12)	19-8-10-1 (15-0)
27/07/2014	Ballymacoda	n = 167	Gill	10 (3/30)	19·8–10·1 (15·0)
Field survey 2					
09/07/2013	Carlingford	n = 5 (T)	Gill	0	20·3–11·9 (16·1)
23/07/2013	Carlingford	n = 5 (T)	Gill	0	20·3–11·9 (16·1)
06/08/2013	Carlingford	n = 5 (T)	Gill	20 (1/5)	18-9-11-15 (15-2)
21/08/2013	Carlingford	n = 5 (T)	Gill	80 (4/5)	18-9-11-15 (15-2)
04/09/2013	Carlingford	n = 5 (T)	Gill	0	16·8-9·1 (13·0)
18/09/2013	Carlingford	n = 5 (T)	Gill	0	16·8-9·1 (13·0)
02/10/2013	Carlingford	n=5 (T)	Gill	20 (1/5)	17:0-11:9 (14:5)
16/10/2013	Carlingford	n=5 (T)	Gill	100 (5/5)	17:0-11:9 (14:5)
30/10/2013	Carlingford	n = 5 (T)	Gill	0	17·0-11·9 (14·5)
22/04/2015	Dungarvan	n = 30 (T)(HS)	Gill	0	14·4-4·2 (9·3)
30/04/2015	Carlingford	n = 30 (T)(HS)	Gill	0	14·2–2·7 (8·5)
06/05/2015	Dungarvan	n = 30 (T)(HS)	Gill	0	14·2-6·7 (10·5)
12/05/2015	Carlingford	n = 30 (T)(HS)	Gill	0	13·3-4·7 (9·0)
18/05/2015	Dungarvan	n = 30 (T)(HS)	Gill	0	14·2-6·7 (10·5)
27/05/2015	Carlingford	n = 30 (T)(HS)	Gill	0	13·3-4·7 (9·0)
03/06/2015	Dungarvan	n = 30 (T)(HS)	Gill	0	16·6–6·2 (11·4)
10/06/2015	Carlingford	n = 30 (T)(HS)	Gill	0	14·3–5·4 (9·9)
16/06/2015	Dungarvan	n = 30 (T)(HS)	Gill	0	16·6–6·2 (11·4)
29/06/2015	Carlingford	n = 30 (T)(HS)	Gill	0	14·3–5·4 (9·9)
01/07/2015	Dungarvan	n = 30 (T)(HS)	Gill	3 (1/30) (T)	19·5–12·1 (15·8)
14/07/2015	Carlingford	n = 30 (T)(HS)	Gill	0	19·0-11·8 (15·8)
16/07/2015	Dungarvan	n = 30 (T)(HS)	Gill	0	19·5–12·1 (15·8)
28/07/2015	Carlingford	n = 30 (T)(HS)	Gill	20 (6/30) (T)	19·0–11·8 (15·8)
30/07/2015	Dungarvan	n = 30 (T)(HS)	Gill	0	19·5–12·1 (15·8)
12/08/2015	Carlingford	n = 30 (T)(HS)	Gill	0	16·4–10·1 (13·3)
13/08/2015	Dungarvan	n = 30 (T)(HS)	Gill	0	18-9-11-5 (15-2)

 $<sup>{}^{</sup>a} Temperature\ data\ after:\ http://www.met.ie/climate/irish-climate-monthly-summary.asp.$ 

August in 2015. Similarly, at Dungarvan Harbour, sampling took place on nine occasions, every 2 weeks from April to August in 2015. All mussel samples were returned to the laboratory and processed immediately for histology and polymerase chain reaction (PCR) analysis to screen for OsHV-1  $\mu$ Var (Table 1).

# Laboratory trials

Naïve Pacific oyster spat (<3 cm, n=600; <1-year-old) were obtained from a hatchery at New Quay, Galway Bay ( $53^{\circ}$  09'16·27"N, 9°04'58·19"W). Oysters from this hatchery have

been used as control oysters in previous studies as the location is deemed uninfected with the herpes virus OsHV-1  $\mu$ Var and variants (Prado-Alvarez *et al.* 2016).

Crassostrea gigas spat were kept in a holding tank (50 L) in a constant temperature (CT) room at  $\sim$ 13 °C in trial 1 and at  $\sim$ 14 °C in trial 2 at a salinity of 35 for 2–3 days until the laboratory cohabitation trials began. The mussel stocks used in the laboratory transmission trials comprised of *Mytilus* spp. from Ballymacoda Bay (trial 1) and *M. edulis* from Carlingford Lough (trial 2).

All seawater used in the laboratory cohabitation trials was sourced from a OsHV-1  $\mu$ Var uninfected site (Bantry Bay, Co.

 $<sup>\</sup>ensuremath{\mathsf{T}}$  – denotes sampling, which occurred at the oyster trestles on the mid intertidal.

HS - denotes sampling, which occurred at the high shore.

Cork, 51°38′59″N, 09°43′00′W) and was UV treated prior to being used.

Laboratory cohabitation transmission trial 1: naïve C. gigas and Mytilus spp. from a OsHV-1  $\mu$ Var endemic culture site held together at 21 °C to determine if the virus can be transmitted between the two species

The combined effects of an elevated seawater temperature (21 °C) on oyster and mussel performance and OsHV-1 μVar pathogenicity were investigated in a laboratory transmission trial. A total of 3×10 L stand-alone control tanks, each containing 40 naïve Galway Bay C. gigas, and three experimental (10L) tanks, each containing 40 naïve Galway Bay C. gigas and 40 randomly sampled wild Mytilus spp. obtained from the Ballymacoda site, which would have experienced a mean seawater temperature in the field of approximately 16 °C were used. An initial sample (n = 30) of oyster and mussels was screened to assess prevalence of the virus at the beginning of the experiment. The control and experimental tanks were placed in a CT room at 21 °C and a salinity of 35. The trial ran for 7 days and the tanks were checked early in the morning and late afternoon for moribund mussels and oysters, which were removed immediately, processed and screened for OsHV-1  $\mu$ Var.

Laboratory cohabitation transmission trial 2: naïve C. gigas and exposed Mytilus spp. from a OsHV-1  $\mu$ Var endemic culture site held under an increasing temperature regimes to determine if transmission could occur between the two species

The combined effects of increasing seawater temperature (14, 21 and 28 °C), at a salinity of 35 on oyster and mussel performance and OsHV-1  $\mu$ Var pathogenicity was investigated in a laboratory trial with wild M. edulis collected from a OsHV-1  $\mu$ Var endemic oyster culture site and hatchery reared naïve C. gigas spat. The M. edulis were randomly collected from in, and around the oyster trestles during the seasonal field sampling conducted at Carlingford Lough in the summer of 2015 (Section Study sites and field sampling). An initial sample (n = 30) of oyster and mussels for both laboratory trials were screened to assess prevalence of the virus at the beginning of the experiment. Two control tanks, each containing 30 naïve Irish hatchery C. gigas, and three experimental tanks, each containing 30 naïve hatchery C. gigas and 30 exposed M. edulis were used. The control tanks and experimental tanks were placed in two separate CT rooms to avoid aerosol contamination. The temperature was initially set in both CT rooms at 14 °C (days 1-14); increased to 21 °C (days 15-20) and subsequently increased to 28 °C (days 21-29) in both CT rooms. The trial ran for 29 days and the tanks were checked early in the morning and late afternoon for moribund or dead oysters and mussels, which were removed and subsequently screened for OsHV-1 µVar. All moribund or dead individuals were screened over the trial and the remaining living oysters and mussels were also screened in both the experimental and control tanks at the end of the trial. A sampling regime of two oysters per tank per day was carried out in the experimental tanks from days 2 to 7. Screening recommenced in both mussels and oysters at day 13 to the end of the trial on day 29 with more randomized screening of approximately six mussels or six oysters being sampled every second day providing an additional 72 living oysters and 71 living mussels from the experimental tanks and 44 living oysters from the control tank being screened. Prior to the commencement of the trial, the shellfish were not cleaned, and over the duration of the trial, the tank water was not changed. The shellfish were fed twice a week using Reed Mariculture Shellfish Diet 1800 (5 ml per tank). Processing of animals from each CT room was carried out in separate laboratories to avoid crosscontamination.

Molecular diagnostic screening of Mytilus spp. and C. gigas DNA extraction and PCR

Mytilus spp. sampled during the first field study (2014) at (a) Ardmore (n = 30) and (b) Garrettstown (n = 60) were screened by PCR for the herpes virus using DNA extracted from gill tissue. From the initial sample (n = 42) collected at (c) Ballymacoda, a subsample of 12 mussels were randomly screened by PCR using gill tissue, haemolymph and shell cavity fluid from each individual. DNA was extracted using the Chelex-100 method (Walsh et al. 1991). All Mytilus spp. sampled during the second field study (2013-2015) at (d) Carlingford Lough and (e) Dungarvan Harbour were screened by PCR for the herpes virus (n = 646)using gill tissue. This screening comprised of 45 Mytilus spp. individuals in 2013 and 601 Mytilus spp. in 2015 with 240 in total from Carlingford lough and 361 from Dungarvan Harbour. Gill tissue (5 mm<sup>2</sup>) was excised and DNA extraction (Qiagen Blood and Tissue kit) was carried out. Mytilus spp. were also screened by PCR for the herpes virus during the laboratory cohabitation transmission trials trial 1 (n = 42) and trial 2 (n = 90).

PCRs using different primer pairs (OHVA/OHVB, OHVC/ OHVD) that detect OsHV-1 and variants (Lynch et al. 2013) were used on all of the mussels sampled from the field sites. Both primer pairs amplify different locations in the C region in the ORF4 gene of the virus (Lynch et al. 2013). All PCRs used a total of 1  $\mu$ L undiluted combined [host + pathogen (if present)] genomic DNA template per individual. The PCR using the OHVA/OHVB primers, which amplify a 385-bp fragment, was carried out in 25 µL L containing 12.9 µL L ddH20, 5 µL, 5× buffer, 5  $\mu$ L dNTPs (0·2 mm), 0·5  $\mu$ L MgCl2 (25 mm stock), 0·25  $\mu$ L of each primer (100 pmol mL<sup>-1</sup> stock) and 0·1  $\mu$ L Taq DNA polymerase. Cycling conditions began with an initial denaturation of the sample at 95 °C for 1 min followed by 35 cycles of 94 °C for 20 s, 56 °C for 30 s and 72 °C for 30 s. A final elongation step took place at 72 °C for 7 min. The PCR using the OHVC/ OHVD primers, which amplify a 296-bp fragment, had the same PCR master mix and thermocycling conditions as described for the OHVA/OHVB PCR (Lynch et al. 2013). Negative controls consisted of deionized distilled water and positive controls were OsHV-1 µVar DNA. DNA was quantified using a spectrophotometer (Thermo-Scientific NanoDrop 1000 spectrophotometer), and extracted DNA was visualized on a 2% agarose gel to ensure DNA was present and to avoid false negatives. Presence of amplified PCR products was confirmed by electrophoresis on a 2% agarose gel stained using 15 mL ethidium bromide (10 mg m $\mathrm{L}^{-1}$ stock) and the gel was run at 110 V for 45 min.

# Quantitative polymerase chain reaction

A subsample of PCR-positive results observed in the mussels from both the second field trial (n = 7) and experimental laboratory cohabitation transmission trials (n = 7) were rescreened by qPCR to detect the viral load (OsHV-1 C region) and mean quantification (range min-max) using HVDP-F and HVDP-R primers (Pépin et al. 2008), which amplify a 197-bp fragment. Negative controls consisting of deionized distilled water were included in the assay. A total of  $5 \mu l$  of  $5 ng mL^{-1}$  combined DNA was screened. The qPCR mix included 12·5 μL qPCR master mix (Agilent Technologies Brilliant SYBR Green QPCR mix) and 12.5 μL each primer. A standard curve using dilutions of five viral genomic DNA suspensions of different quantities  $(1\times10^5, 1\times10^4, 1\times10^3, 1\times10^2, 1\times10)$  in triplicate was used. Amplification included a denaturation step of 10 min at 95 °C and 40 amplification cycles (30 s at 95 °C, 1 min at 60 °C and 45 s at 72 °C). Melting temperature curve analysis was 1 min at 95 °C, 30 s at 60 °C and 30 s at 95 °C. All samples were tested in duplicate (Lynch et al. 2013). Viral load was classified into

three categories as described by Renault *et al.* (2014): (i) category 0, no viral DNA; (ii) category 1, viral-DNA amounts of <104 DNA copies per mg of wet tissue; and (iii) category 2, viral-DNA amounts of >104 DNA copies per mg of wet tissue.

#### Sequencina

Direct Sanger sequencing on PCR products (385-bp) amplified from the samples using the OHVA/OHVB primers was carried out to ensure the amplified sequences were being recovered from the OsHV-1 genome. DNA isolated from PCR products amplified from separate mussel individuals (n=5) from each site were pooled into replicates from each individual (4–5) to increase the DNA concentration. Both the forward and reverse strands of DNA samples were sequenced commercially (Eurofins MWG Operon, Ebersberg, Germany). Each sequence was matched against a nucleotide database (https://blast.ncbi.nlm.nih.gov/) to confirm the individual sequences were from the OsHV-1 and variants genome.

# Histological screening

A subsample of mussels (n = 20) that were collected at Carlingford Lough in 2013 were prepared for histological analysis and in situ hybridization (ISH). For each animal, a 5 mm crosssection section of digestive gland, gonad, gills and mantle was removed and a 3 mm cross-section of the visceral mass was excised in front of the pericardial region for histological analysis and immediately fixed in Davidson's solution at 4 °C for 48 h after which they were placed in 70% ethanol. Samples were processed (Shandon Citadel 1000) and microtomed, 5 µM tissue sections for histology and 7 µM for ISH. Tissue sections were stained with haematoxylin and eosin while a digoxygenin (DIG) labelled probe was used for ISH. Sections were viewed with a Nikon Eclipse 80i and images were captured using NIS elements software (at  $100\times$ ,  $200\times$ ,  $400\times$  and  $1000\times$ . The presence of any abnormal and/or viral cells (Renault et al. 1994a; Renault and Novoa, 2004) was noted.

# ISH (DIG labelled probe)

ISH analysis was conducted in accordance with the Ifremer protocol (Renault and Lipart, 1998; Lipart and Renault, 2002; Segarra *et al.* 2016); however, the OHVA/B and OHVC/D primer pairs were used to create the probe (Lynch *et al.* 2013). ISH was conducted on the 20 Carlingford samples on silane-prep<sup>™</sup> slides. A DIG labelled probe was produced using OsHV-1 primers and OsHV-1 DNA as a template which was used to hybridize to the target viral DNA within the tissue samples. The slides were mounted using Eukiit mounting medium and cover slips. As with histology, slides were viewed using a microscope (100×, 200×, 400× and 1000×) (Nikon Eclipse 80i) and images were captured using NIS elements software.

#### Data analysis

Statistical analyses were carried out using the IBM SPSS Statistics version 23. Statistical significance was determined using P < 0.05. Yates's continuity corrected  $\chi^2$  test was performed to compare percentage (%) mortality between experimental and control groups. Fisher's exact test was used to compare percentage (%) infected between experimental and control groups and to compare percentage (%) mortality in infected and non-infected animals. Fisher's exact test was used when one or more cells had an expected count of <5.

#### **Results**

## Seasonal field sampling

## Field survey 1

In the preliminary field survey conducted in 2014, a prevalence of infection of 7% (2/30) was observed in the initial sample at (a) Ardmore and 3-3% (1/30) in (b) Garrettstown. In the second Garrettstown mussel sample taken after mussels were placed into holding tanks for 3 weeks in the laboratory at 13 °C, a significant increase in virus detection with 46-7% (14/30) prevalence was observed. At (c) Ballymacoda, all mussels (n = 12) screened were positive for OsHV-1  $\mu$ Var in each sample type, gill tissue, haemolymph and shell cavity fluid, taken from each mussel (Table 2).

## Field survey 2

OsHV-1  $\mu$ Var DNA was detected in *Mytilus* spp. at (c) Carlingford Lough in 24% of the mussels screened at the oyster trestles (11/45) in 2013 and 2% (6/240) in 2015. The six Carlingford Lough mussels had a category 1 viral load with a range of  $2\cdot22\times10^2$  to  $1\cdot565\times10^3$  viral copies  $\mu$ L<sup>-1</sup> of genomic DNA (n=6) with a mean value of  $6\cdot8\times10^2$  viral copies  $\mu$ L<sup>-1</sup> of genomic DNA. In the 2015 field study, prevalence of infection in *C. gigas* seed were minimal at Carlingford Lough at 10% (25/240). During the second field trial (2013–2015), a single mussel (0·2%, n=361) was positive for OsHV-1  $\mu$ Var and had a category 1 viral load with  $1\cdot23\times10^2$  viral copies  $\mu$ L<sup>-1</sup> of genomic DNA at the trestles at (e) Dungarvan Harbour. qPCR was not conducted on the 2013 samples due to the degradation and reduction in quality of the genomic DNA (Table 2).

Laboratory cohabitation transmission trial 1: naïve C. gigas held with Mytilus spp. collected 400 m from a OsHV-1  $\mu$ Var endemic culture site

In laboratory trial 1, the initial sample (n = 30) of Galway *C. gigas* was negative for OsHV-1  $\mu$ Var; however, the initial sample (n = 12) of *Mytilus* spp. from Ballymacoda was positive for the virus with 100% (12/12) prevalence, using representative DNA from

**Table 2.** Summary of all mussel samples screened for OsHV-1  $\mu$ Var at the five Irish sites in field surveys 1 and 2 in 2013–2015

Sample year	Sample site	PCR (% prevalence)	Infection intensity DNA copies per mg of wet tissue <sup>a</sup>
2013	Carlingford Lough	24% (11/45)	No qPCR conducted
2014	Ardmore	7% (2/30)	No qPCR conducted
2014	Garretstown	25% (15/60)	No qPCR conducted
2014	Ballymacoda	100% (12/12)	No qPCR conducted
2014	Ballymacoda	10% (3/30)	No qPCR conducted
2015	Carlingford Lough	2% (6/240)	Category 1 < 10 <sup>4</sup>
2015	Dungarvan Harbour	0.2% (1/361)	Category 1 < 10 <sup>4</sup>

<sup>&</sup>lt;sup>a</sup>Infection intensity after Renault *et al.* (2014).

(i) gill tissue (n = 12), (ii) haemolymph (n = 12) and (iii) the shell cavity fluid (n = 12). No oyster or mussel mortalities were observed in the 7-day trial. During the trial, OsHV-1  $\mu$ Var was not detected in the control oysters 0% (0/15); however, 10% (3/30) of the experimental mussels were positive and 100% (30/30) of the experimental oysters (i.e. naïve oysters exposed to infected mussels) were positive for OsHV-1  $\mu$ Var.

Laboratory cohabitation transmission trial 2: naïve C. gigas and exposed Mytilus spp. collected at oyster trestles held at various increasing temperatures over a 4-week period in 2015

In laboratory trial 2, mortality was 26·7% (16/60) in the control group (oysters) and 10% (18/180) in the experimental groups [9% (8/90) in the experimental oysters and 11% (10/90) in the experimental mussels] (Fig. 1). A total of 14·1% (34/240) of the total animals tested, died over the duration of the study. Yate's continuity corrected  $\chi^2$  test indicated that the percentage (%) mortality differed significantly between the control and experimental groups [ $\chi^2$  (1) = 8·96, P<0·0·1]; experimental group had lower % mortality. Of the 16 (26·7%) oysters who died in the control groups, all were being held at 28 °C. Of the 18 (20%) animals (mussels and oysters) that died in the experimental groups, three mortalities occurred in the oysters at 14 °C and, five oyster and 10 mussel mortalities occurred at 28 °C (Fig. 1).

The initial samples (n = 30) of Galway *C. gigas* and Carlingford Lough M. edulis were both negative for OsHV-1 μVar. In the experimental groups, 3.9% (7/180) of animals [mussels (n = 2) and oysters (n = 5)] were positive for OsHV-1  $\mu$ Var compared with 3.3% (2/60) in the control oyster group (Fig. 1). Fisher's exact test indicated that there was no significant difference in prevalence (%) of the virus between treatments (experimental and control groups) (P > 0.05). Of the seven infected animals in the treatment groups, a single oyster was infected at 14 °C (alive at the end of the study), and the remaining six individuals (four oysters and two mussels) were infected at 28 °C (five were alive and one mussel was dead at the end of the study). The two infected control animals occurred at 28 °C were found moribund at the end of the study. Of the positive oysters (n = 5) and mussels (n = 2), higher viral loads were observed in the oysters  $(n = 5, 3.84 \times 10^2 \text{ to } 3.67 \times 10^3 \text{ viral copies } \mu\text{L}^{-1} \text{ of genomic DNA})$  with an average of  $1.08 \times 10^3 \text{ viral copies } \mu\text{L}^{-1} \text{ of genomic DNA})$  than the mussels  $(n=2, 4\cdot16\times10^2 \text{ to } 6\cdot6\times10^2 \text{ viral copies }\mu\text{L}^{-1}$  of genomic DNA with an average of  $5\cdot38\times10^2$  viral copies  $\mu\text{L}^{-1}$  of genomic DNA) (Table 3). These are classified as category 1, viral-DNA amounts of  $<10^4$  DNA copies per mg of wet tissue (Renault *et al.* 2014). Fisher's exact test revealed that there was no statistically significant difference in percentage (%) mortality between individuals with the virus and those without (oysters and mussels) (P>0.05). Mortality rates were  $11\cdot1\%$  (1/9) in the infected group compared with  $14\cdot3\%$  (33/231) in the non-infected group.

## Histological examination and ISH

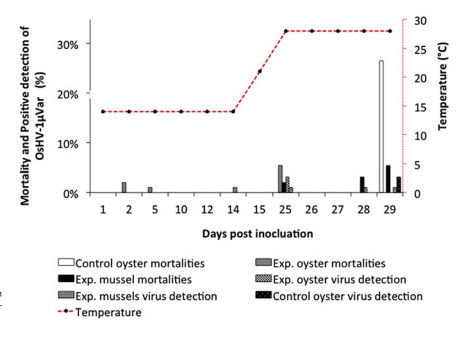
Abnormal cells consistent with the presence of OsHV-1  $\mu$ Var were observed in the *Mytilus* spp. histology in individuals that were positive for OsHV-1  $\mu$ Var by PCR. In *Mytilus* spp., collected from the field (Ballymacoda), abnormal cells, which were enlarged, rounded with the nuclei disrupted or pushed to the edge of the cell were detected in half (10/20) of the histology samples screened and were mostly observed throughout the connective tissue in the mantle and to a lesser extent in the gills and digestive tubules (Fig. 2). In the ISH analysis, a signal was not detected in individuals deemed to be uninfected with OsHV-1  $\mu$ Var in the PCR (Fig. 3), while a positive signal for OsHV-1  $\mu$ Var in 20% (4/20) of the mussels screened was detected using this method (Fig. 4).

# Sequencing

Five forward and reverse sequences from five individual mussels, which tested positive in the PCR, were generated in the direct Sanger sequencing. All forward and reverse sequences were identical and all sequences were confirmed by Blastn to be from the OsHV-1 genome with a 100% maximum identity and 99% query coverage (GenBank accession no. KU861511·1) with OsHV-1 strain  $\mu$ Var isolate Cg\_C2C6 region genomic sequence.

# Discussion

In this study, OsHV-1  $\mu$ Var was detected in wild *Mytilus* spp. and *M. edulis* tissues with associated pathologies similar to those previously reported in oysters infected by herpes virus (Renault *et al.* 



**Fig. 1.** Mortality (%) and prevalence (%) of OsHV-1  $\mu$ Var in co-habiting naïve *Crassostrea gigas* and *Mytilus* spp., naturally exposed to OsHV-1 $\mu$ Var, in laboratory trial 2.

Table 3. Summary of Crassostrea gigas and Mytilus spp. screened for OsHV-1 μVar in the laboratory cohabitation transmission trial 2 in 2015

Sample type	Virus prevalence (%)	Infection intensity DNA copies per mg of wet tissue <sup>a</sup>	Mortality (%)
TRIAl 2_Initial <i>C. gigas</i>	0% (0/30)	Not screened in qPCR	N/A
TRIAl 2_Initial <i>Mytilus</i> spp.	0% (0/30)	Not screened in qPCR	N/A
TRIAL 2_Experimental Mytilus spp.	2% (2/90)	Category 1 < 10 <sup>4</sup>	9% (8/90)
TRIAL 2_Experimental <i>C. gigas</i>	6% (5/90)	Category 1<10 <sup>4</sup>	11% (10/90)
TRIAl 2_CONTROL <i>C. gigas</i>	3% (2/60)	Not screened in qPCR	27% (16/60)

<sup>&</sup>lt;sup>a</sup>Infection intensity after Renault et al. (2014).

1994a; Renault and Novoa, 2004). Although these histological changes and abnormalities associated with OsHV have not been previously described for Mytilus spp., similar changes have formerly been observed in a non-ostreid bivalve species, the Manila clam R. philippinarum (Renault et al. 2001). Of significance, this study reports the first transmission of this virus from both M. edulis and Mytilus spp. to naïve C. gigas, which occurred in the laboratory under an increased and increasing temperature regime; however, it must also be highlighted that replication of the virus with a 14-fold increase also occurred in mussels, which were held in the laboratory at 13 °C during a 3-week period. The virus was detected in mussels cohabiting with oysters within the oyster bags at the trestles but it was also detected in wild mussels located 400 and 500 m from the oyster trestles and up to 26 km from the oyster farms at non-culture sites. The impact of the virus on mussel health is unclear, as OsHV-1  $\mu$ Var was detected at an overall low prevalence in mussels and at a low viral load for those screened by qPCR and without associated mortality in the laboratory trials; however, cytopathic effects were observed in the mussels similar to those observed in C. gigas, which have been associated with oyster mortalities. A low prevalence of OsHV-1 µVar was observed in oyster populations in the field at the same time, possibly due to lower than average seawater and air temperatures experienced at those sites for the duration of the trials.

According to the World Organization for Animal Health (OIE), a 'suspected' case of infection with microvariants, can exist without evidence of mortality, whereas a 'confirmed' case

occurs when detection by histology, transmission electron microscopy or PCR is followed by sequencing confirmation. In this study, detection of OsHV-1 µVar DNA was established by PCR in mussel samples from all five study sites, in the qPCR screening of mussel samples from two of the study sites, and visualized by histology and ISH in mussels from a single site. The DNA was confirmed by direct sequencing to be that of OsHV-1 µVar in samples from each study site. From the qPCR analysis, it can be concluded that both field sampled and laboratory experimental living and dead Mytilus spp. had detectable quantities of OsHV-1  $\mu$ Var. The mussels used in the laboratory cohabitation transmission trial 2 were exclusively from Carlingford Lough, which represents the first record of OsHV-1  $\mu$ Var in M. edulis. In contrast to our study, OsHV-1 DNA was previously detected in samples of *Mytilus* spp. in Australia with a 20% (n = 1/5) prevalence and viral concentrations below the quantification limit of the qPCR assay (<12 DNA copies per PCR reaction) (Evans et al. 2017); however, this was not the case in this study where up to 1565-37 viral copies  $\mu$ L<sup>-1</sup> of genomic DNA were detected in Irish mussels. In addition, abnormal cells consistent with the pathology associated with OsHV-1  $\mu$ Var infection were visualized in the mussel histology. The cytopathic effects (abnormal cell morphology) observed in infected mussels were similar to those visualized in infected oysters in previous studies (Renault et al. 1994a, 2000; Renault and Novoa, 2004). This study demonstrated that elevated temperatures exacerbated OsHV-1  $\mu$ Var prevalence and transmission between mussels and oysters in the laboratory but mortality was relatively low in both species. The

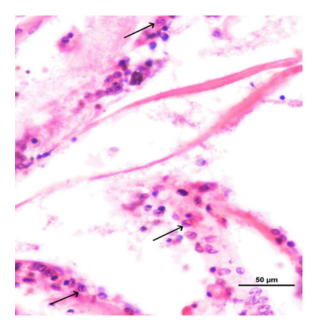
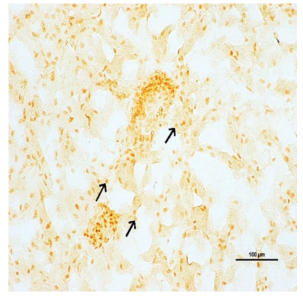
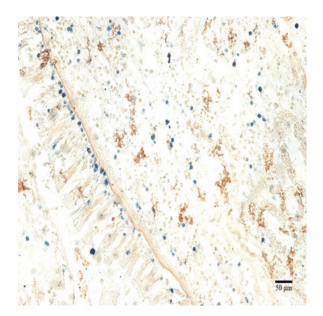


Fig. 2. Tissue section of Mytilus spp. with OsHV-1  $\mu$ Var-infected blood cells.



**Fig. 3.** *In situ* hybridization stained *Mytilus* spp. connective tissue of normal cells (orange-brown staining indicates uninfected, healthy normal tissue) (arrowheads).



**Fig. 4.** *In situ* hybridization stained *Mytilus* spp. digestive epithelia tissue (left of image) and connective tissue showing cells infected with OsHV-1 microVar (stained blue) (orange-brown staining indicates uninfected, healthy tissue).

differences in oyster performance in the laboratory, be they control or experimental animals, could be attributed to natural variation in the immune function and performance of individual oysters even within the same population/stock. During the summer of 2013 air temperatures higher than average (up to 3.5 °C above the average temperature) were experienced (Prado-Alvarez *et al.* 2016), while in the summer of 2015 all monthly mean air temperatures were below their long-term average (www.met.ie) and positive detection of the virus was observed primarily in July, August and October during phases of warmer weather and dry periods with low precipitation.

OsHV-1  $\mu$ Var is generally detected in dying oysters when seawater temperatures are >16 °C (Pernet et al. 2012). Higher temperatures appear to act as a stressor for C. gigas and exacerbate virus proliferation and transmission; and detection and mortality are associated with a marked increase in mean daily seawater temperature (Garcia et al. 2011). Previous studies of Mytilus spp. have also observed that temperature is also a key driver of biological response (Hu et al. 2015) with elevated temperatures negatively impacting on the immune response in M. edulis by significantly increasing the antibacterial activity of cell-free haemolymph (Ellis et al. 2015). Crassostrea gigas has a broad temperature tolerance, with a range of -1.8 to 35 °C (FAO, 2015); however, higher temperatures exacerbate virus proliferation, transmission, and mortality as was observed in this present study. OsHV-1  $\mu$ Var is known to be persistent in oysters held at temperatures as low as 13 °C and reactivated during thermal elevation to 21 °C (Pernet et al. 2015), this was confirmed by the findings of our study following elevation of temperature during the laboratory trial. However, our results would further indicate that the virus is replicating also at 13 °C in the mussels. The stressful conditions of the artificial laboratory holding conditions and the close proximity of the mussels to one another in the tank/s for a prolonged period of time may have facilitated the activation of the virus in the mussels and its subsequent replication within a system where no water changes occurred. It may also be possible that the virus responds differently to the virus at a reduced temperature in mussels compared with oysters.

OsHV-1  $\mu$ Var was detected in a small percentage of the control living oysters; however, this may have been attributed to background aerosol contamination as the initial sample was negative

and this population is classified as disease free. Elevated temperature alone may not result in increased mortalities, e.g. C. gigas that experienced prolonged high temperature (21 °C for 14 days) in the Inland Sea, Wales, did not experience mass mortalities (Malham et al. 2009). However, in the context of a changing climate, with future increases in seawater temperatures predicted, the ecological sustainability of marine fisheries and aquaculture may be threatened (Dang et al. 2012). Consideration of these results should be given in areas where both mussel and oyster cultures occur and for the movement of mussels from OsHV-1  $\mu$ Var infected sites to sites where Pacific oysters may be cultured and are free of infection. Even with these viral loads and low number of animals testing positive, transmission of the virus was still effected to naive oysters, which highlights the risks involved with the movement of shellfish for aquaculture and the unintentional introduction of pathogens in non-typical hosts. Wild mussels approximately 400 m from a C. gigas culture site successfully transmitted OsHV-1  $\mu$ Var to naïve *C. gigas* in the laboratory, thus indicating that the virus is viable and is being maintained outside the known host. Of interest, the virus was detected in all of the experimental naïve oysters that were held with exposed mussels in laboratory trial 1.

The full process of how the virus might maintain itself at Pacific oyster culture sites is unclear and the effects of OsHV-1  $\mu$ Var infection on the mussel population in Ireland, in particular those at infected oyster culture sites, is yet to be determined. *Mytilus* spp. appears to act as a reservoir or carrier for OsHV-1  $\mu$ Var but the nature of this interaction and impacts on mussel health need to be investigated further. What is certain from the findings of this study is that OsHV-1  $\mu$ Var has the ability to be associated with other cohabiting bivalve species even at a distance from a disease 'hot spot' such as an OsHV-1  $\mu$ Var endemic *C. gigas* culture site.

**Acknowledgements.** The authors would like to acknowledge and thank the shellfish growers for their assistance during fieldwork and for provision of samples.

**Financial support.** This study was carried out with financial assistance from the EUFP7 Bivalife Project 'Controlling infectious diseases in oysters and mussels in Europe'. Project number: 266157. Call (part) identifier: FP7-KBBE-2010-4 Improving European mollusc aquaculture: disease detection and management. This study was also carried out with financial assistance from the REPOSUS Project 'Reducing the impact of Pathogens and disease in the Irish Oyster industry to support the sustainability and growth of the sector' funded under FIRM by DAFM 14 SF 820.

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