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# Diversity and molecular epidemiology of *Ostreid herpesvirus 1* in farmed *Crassostrea gigas* in Australia: Geographic clusters and implications for "microvariants" in global mortality events

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#### ABSTRACT

Since 2010, mass mortality events known as Pacific oyster mortality syndrome (POMS) have occurred in Crassostrea gigas in Australia associated with Ostreid herpesvirus 1. The virus was thought to be an OsHV-1 uVar or "microvariant", i.e. one of the dominant variants associated with POMS in Europe, but there are few data to characterize the genotype in Australia. Consequently, the genetic identity and diversity of the virus was determined to understand the epidemiology of the disease in Australia. Samples were analysed from diseased C. gigas over five summer seasons between 2011 and 2016 in POMS-affected estuaries: Georges River in New South Wales (NSW), Hawkesbury River (NSW) and Pitt Water in Tasmania. Sequencing was attempted for six genomic regions. Numerous variants were identified among these regions (n = 100 isolates) while twelve variants were identified from concatenated nucleotide sequences (n = 61 isolates). Nucleotide diversity of the seven genotypes of C region among Australian isolates (Pi 0.99  $\times$  10<sup>-3</sup>) was the lowest globally. All Australian isolates grouped in a cluster distinct from other OsHV-1 isolates worldwide. This is the first report that Australian outbreaks of POMS were associated with OsHV-1 distinct from OsHV-1 reference genotype, µVar and other microvariants from other countries. The findings illustrate that microvariants are not the only variants of OsHV-1 associated with mass mortality events in C. gigas. In addition, there was mutually exclusive spatial clustering of viral genomic and amino acid sequence variants between estuaries, and a possible association between genotype/amino acid sequence and the prevalence and severity of POMS, as this differed between these estuaries. The sequencing findings supported prior epidemiological evidence for environmental reservoirs of OsHV-1 for POMS outbreaks in Australia.

#### 1. Introduction

Since late last century, the Pacific oyster, *Crassostrea gigas*, has been subject to recurrent disease events (Nicolas et al., 1992; Renault et al., 2000; Renault et al., 1994; Samain et al., 2007). However, in 2008 a marked increase in mass mortality outbreaks occurred in France (Cochennec-Laureau et al., 2010; Martenot et al., 2011; Segarra et al., 2010) and later in other areas in the world (Abbadi et al., 2018; Batista et al., 2015; Gittenberger et al., 2016; Hwang et al., 2013; Keeling et al.,

2014; Mortensen et al., 2016; Peeler et al., 2012; Roque et al., 2012; Segarra et al., 2010). The disease was called Pacific oyster mortality syndrome (POMS) when it first appeared in Australia in 2010 in the Georges and Parramatta rivers, which are both major estuaries in Sydney (Jenkins et al., 2013; Paul-Pont et al., 2013a). POMS causes considerable economic losses in Australia, New Zealand and Europe (Fuhrmann et al., 2019). It is a multifactorial disease influenced by environmental conditions such as water temperature and host characteristics such as age and size (Hick et al., 2018; Petton et al., 2013) and

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although it has a proposed polymicrobial etiology involving bacteria (de Lorgeril et al., 2018; Friedman et al., 2005; Petton et al., 2021), outbreaks are induced by infection with *Ostreid herpesvirus 1* microvariants (Batista et al., 2015; Gittenberger et al., 2016; Hwang et al., 2013; Jenkins et al., 2013; Keeling et al., 2014; Mortensen et al., 2016; Peeler et al., 2012; Renault et al., 2012; Roque et al., 2012). The disease typically occurs in summer and mortality can be reproduced experimentally by controlled exposure of oysters to the virus alone in a dose-responsive and temperature-dependent manner (de Kantzow et al., 2016; Paul-Pont et al., 2015).

Ostreid herpesvirus 1, OsHV-1, which is the only species included in the genus Ostreavirus, is a double-stranded DNA virus (Davison et al., 2009). The presence of the gene coding for the ATPase subunit of a terminase involved in the packaging of viral DNA inside the capsid, in addition to intranuclear replication, places OsHV-1 in the Malacoherpesviridae under the order Herpesvirales (Davison et al., 2009; Davison et al., 2005). Members of this taxon are widespread, both in terms of the range of species infected and also geographically. They have been detected in many hosts besides C. gigas including Crassostrea virginica (Farley et al., 1972), Ostrea angasi (Hine and Thorne, 1997), Ostrea edulis (Comps and Cochennec, 1993), Tiostrea chilensis (Hine et al., 1998), Pecten maximus (Arzul et al., 2001a), Ruditapes philippinarum (Renault et al., 2001), Ruditapes decusstaus (Renault and Arzul, 2001), Crassostrea angulata (Batista et al., 2015), Scapharca broughtonii (Bai et al., 2016; Xia et al., 2015), Carcinus maenas (Bookelaar et al., 2018), Chlamys farreri (Bai et al., 2019), Octopus vulgaris (Prado-Alvarez et al., 2021) and in numerous sites around the world including Europe, Asia, Oceania and the Americas (Abbadi et al., 2018; Batista et al., 2015; Burge et al., 2021; Gittenberger et al., 2016; Hwang et al., 2013; Jenkins et al., 2013; Keeling et al., 2014; Mortensen et al., 2016; Peeler et al., 2012; Roque et al., 2012; Segarra et al., 2010; Shimahara et al., 2012).

A specific microvariant, OsHV-1 µVar, was described in association with the large increase in mortality in C. gigas on the French coast in 2008 (Segarra et al., 2010), but the detection of this virus in a specimen collected before 2008 (Martenot et al., 2012) raises the question of its actual or exclusive involvement in the increasing global phenomenon of oyster mortality. OsHV-1 µVar was strictly characterised by mutations upstream of ORF 4 consisting mainly of 12 deletions in the microsatellite zone and a few mutations upstream and downstream which impact the amino acid sequence. In addition, at ORF 42-43 OsHV-1 µVar had a subtitution of a thymine instead of a cytosine downstream of the sequence without showing any impact on the amino acid sequence (Segarra et al., 2010). Following the publication of the OsHV-1µVar sequence from region C, partly located at ORF 4 (Segarra et al., 2010), several isolates were analysed and this revealed a range of « microvariants » close to the OsHV-1 µVar sequence (Burge et al., 2021; Burioli et al., 2018; Martenot et al., 2012; Morga et al., 2021). Data for numerous OsHV-1 genotypes characterised by sequencing of various coding and non-coding regions or whole genomes are available: in Europe, the reference genome isolated from C. gigas larvae (Davison et al., 2005), the variant OsHV-1var sampled from P. maximus (Arzul et al., 2001a) and the microvariant OsHV-1 µVar from C. gigas spat and juveniles (Abbadi et al., 2018; Burioli et al., 2017; Segarra et al., 2010); in China, Acute Viral Necrosis Virus (Ren et al., 2013), OsHV-1-SB isolated from S. broughtonii (Bai et al., 2016; Xia et al., 2015) and the variant isolated from C. farreri (Bai et al., 2019); in Sweden OsHV-1 SW6 found in O. edulis (Morga et al., 2021); and in many parts of the world, so called « microvariants » or  $\mu$ Vars, which are similar to OsHV-1  $\mu$ Var (Burge et al., 2021; Morga et al., 2021). Double-stranded DNA viruses are known to be genetically stable; DNA viruses in general have mutation rates of  $10^{-8}$  to  $10^{-6}$  substitutions per nucleotide site per cell infection, a lower rate than RNA viruses, due the viral DNA polymerase that exhibits proofreading activity (Peck and Lauring, 2018). However, Morga et al. (2021) recently reported a faster evolutionary rate than usual for OsHV-1. The emergence of variations may be due to selection pressure and the ability of the virus to adapt to its environment and host as reported in other members of the *Herpesvirales* (Renzette et al., 2013). The available data suggest that pathogenic subtypes of OsHV-1 known as microvariants possess genetic diversity beyond that defined for a single genotype by Segarra et al. (2010), but the extent of diversity and how this relates to their global emergence is unclear.

The first detection of OsHV-1 associated with mortality of *C. gigas* in Australia occured in 2010 but despite recurrent outbreaks there has been little characterisation of the genetic diversity of the viruses affecting *C. gigas* in Australia (Jenkins et al., 2013). Available data suggest that the genotype responsible for the index case in 2010 was close to OsHV-1  $\mu$ Var but had some differences (Jenkins et al., 2013). Given that OsHV-1 genetic diversity could affect the gravity of the disease (Delmotte et al., 2022; Martenot et al., 2011; Morga et al., 2021), it is important to study the genetic composition of the Australian isolates to better understand epidemiological patterns in Australia. Detailed, long term, epidemiological studies in Australia revealed that POMS outbreaks recurred at the same sites over successive years (Whittington et al., 2019). The source of OsHV-1 for these disease events between 2011 and 2016 was environmental rather than spread through oyster farming activities (Whittington et al., 2018).

The aim of this report is to (i) study the identity and genetic diversity of OsHV-1 in archival samples collected during mortality events in Australia using six regions sequenced by Sanger technology, (ii) compare the Australian viruses with international isolates previously published, and iii) make inferences based on diversity and phylogeny about the source and transmission of the virus to inform management decisions for disease prevention and control.

#### 2. Materials and methods

#### 2.1. Oyster sample selection

Oyster samples were selected over the five summer seasons between 2011 and 2016 from the entire geographical distribution of POMS disease outbreaks in commercial aquaculture in Australia. Comprehensive data for each sample including location, age, size, mortality rate and viral load are provided in Supplementary data 1, linked by references to scientific publications on these populations. In summary, all samples were collected on oyster farms by researchers from the University of Sydney and all oysters in the final dataset were initially recruited as spat from an OsHV-1 free hatchery in Tasmania (Table 1). All samples were tested using OsHV-1 real-time PCR (Martenot et al., 2010) and positive samples containing the highest concentration of OsHV-1 DNA from among the samples collected that day were selected for sequencing. A total of 118 samples were selected and sequence was subsequently obtained from at least one region of the genome from 107 samples to which a unique "isolate ID" was then assigned. For seven of these (all wild C. gigas or Saccostrea commercialis from the Georges River in the 2013-2014 season) sequence was obtained only for Region 3 which had no variation (see below); these isolates were excluded from further analysis, leaving 100 isolates available for analysis, all being samples from farmed C. gigas (Supplementary data 1). Of the 100 samples of C. gigas, 38 were collected during periods of OsHV-1 infection in the

Table 1

Number of samples of OsHV-1 PCR-positive oysters from which nucleotide sequences were analysed according to the estuary and the season of collection. The number of sites sampled are in parentheses; sites differed between seasons.

Estuary	2011- 2012	2012- 2013	2013- 2014	2014- 2015	2015- 2016	Total
Georges River NSW	19 (3)	9 (3)	14 (4)	13 (3)	5 (3)	60
Hawkesbury River NSW		11 (3)	15 (2)	9 (2)	3 (2)	38
Pitt Water TAS Total	19	20	29	22	2 (1) 10	2 100

Hawkesbury River estuary north of Sydney NSW (from Coba Bay n = 2; Kimmerikong Bay n = 5; Mooney Mooney n = 15; Mullet Creek n = 10; Porto Bay n = 6), 60 were collected in the Georges River estuary south of Sydney NSW (from Limekiln Bar n = 1; Neverfail Bay n = 2; Site A n =13; Site B n = 10; Site C n = 25; Sylvania Waters n = 1; The Shed n = 8), while two were collected in Tasmania (from Pitt Water n = 2) (Fig. 1). Exact locations are illustrated in maps in Whittington et al. (2019) and in de Kantzow et al. (2017). Sampled oysters had an age range of 5.5 months to 17 months and a size range of 5mm to 110mm. They were all the subject of published studies in which details of the oyster populations are reported (de Kantzow et al., 2017; Hick et al., 2018; Paul-Pont et al., 2013a; Paul-Pont et al., 2013b; Paul-Pont et al., 2014; Whittington et al., 2015a; Whittington et al., 2015b; Whittington et al., 2019) (Supplementary data 1).

#### 2.2. Description of the selected regions

Six regions of the OsHV-1 genome were targeted for Sanger sequencing. Firstly, four regions were selected based on their discriminatory value for distinguishing microvariant genotypes from among other OsHV-1 variants. Region 1 and Region 2 targeted ORF 4 and ORF 43, respectively. They were selected in order to distinguish OsHV-1µVar (Segarra et al., 2010), OsHV-1µVar∆9, OsHV-1µVar∆ 15 described in Europe (Martenot et al., 2012; Martenot et al., 2011) and other microvariants (Bai et al., 2016). Region 3 covered ORFs 35, 36, 37 and 38 and bounded a large deletion in OsHV-1µVar that induced the suppression of ORF 36 and 37 and disrupted ORF 38 (Burioli et al., 2017; Martenot et al., 2015; Martenot et al., 2013; Morga et al., 2021; Renault et al., 2012). Region 4 was selected for the high level of polymorphism suitable for distinguishing genotypes of OsHV-1 (Batista et al., 2015). It is located between ORF 49 and ORF 50 and is a non-coding site. Region 5 located on ORF 11, showed a large deletion of 1386bp in microvariants (Morga et al., 2021; Burioli et al., 2017; Martenot et al., 2013). Finally, Region 6, which could be involved in virulence mechanisms is a polymorphic region of interest; it targets ORF 88 coding for a transmembrane glycoprotein (Martenot et al., 2013). PCR and sequencing reaction primers used for each region are provided in Table 2.

#### 2.3. Sample processing

#### 2.3.1. Extraction

Each sample was derived from whole animal or dissected mantle and gill tissues of one or more oysters, depending on their size (Supplementary data 1). The DNA extraction was performed according to previous studies (Evans et al., 2014; Whittington et al., 2019). Briefly, oysters were collected in the field and transported directly to the laboratory where they were frozen at  $-80^{\circ}$ C until processed. A  $400\pm100$  mg sample was homogenised by bead beating using sterile stainless steel beads (Aussie Sapphires) and 1 mL distilled water and placed into a TissueLyser II machine (Qiagen) for 2 min at frequency 30, repeated once. All samples were clarified by centrifugation at 1340 g for 2 min in a microcentrifuge and supernatants were removed and stored in sterile tubes at -80°C. Nucleic acids were purified using a 5X MagMAXTM-96Viral RNA Isolation Kit (Ambion, Life TechnologiesTM, Mulgrave, Australia) and then MagMAXTM Express 96 magnetic particle processor (Applied BiosystemsTM, Life TechnologiesTM, Mulgrave, Australia) according to manufacturers' instructions for a final volume of 50 µL using the AM1836 deep-well standard programme (Ambion, Life TechnologiesTM, Mulgrave, Australia). Purified nucleic acids were stored at  $-20^{\circ}$ C.

#### 2.3.2. PCR

Various PCR were perfomed in order to target the 6 regions described above in part 2.2. The primers are reported in Table 2. For Region 1 and Region 2, each reaction contained 5  $\mu$ l DNA extract, 5 $\mu$ l 10X PCR buffer (66.6mM Tris-HCl,16.6mM (NH4)2SO4, 2.5mM MgCl2, 1.65mg/ml bovine serum albumin,10mM beta-mercaptoethanol), 10 $\mu$ l dNTP mix (1mM), 250nM each forward and reverse primer, 5U DNA polymerase (taq:pfu mix) and nuclease-free water to a final volume of 50 $\mu$ l. Thermocycling was performed (Corbett Research CGI960) according to the following conditions: 1 cycle at 94°C for 2min; 35 cycles consisted in 94°C for 1min, 50°C for 1 min and 72°C for 1min, and a final extension



Fig. 1. Sampling sites. George river estuary sites are located on the bottom map corresponding to the south of Sydney and Hawkesbury river estuary sites are located on the upper map corresponding to the north of Sydney.

Regions of the OsHV-1 genome targeted for PCR amplification.

Sequencing	Amplificatio		Refs.			
region	Position Gene		Primer name and purpose <sup>b</sup>	Primer sequence $(5' \rightarrow 3')$	Amplicon size (base pairs)	
1	178211	ORF 4	C2 (A, S)	CTCTTTACCATGAAGATACCCACC	709	(Renault and Arzul,
(C region)	178919		C6 (A, S)	GTGCACGGCTTACCATTTTT		2001)
			C1585 (S)	GTATAAATAGGCGCG		This paper
			C1586 (S)	CGCGCCTATTTATAC		This paper
			C1587 (S)	CAGACGAGGTTAAC		This paper
			C1588 (S)	GTTAACCTCGTCTG		This paper
			C1593 (S)	GGAGCTGCGGCGCTATG		This paper
2	59950	ORF 42-43	IA1 (A, S)	CGCGGTTCATATCCAAAGTT	607	Segarra et al. (2010)
(A region)	60557		IA2 (A, S)	AATCCCCATGTTTCTTGCTG		
			C1589 (S)	CTTGCTCATCGTATTC		This paper
3	51979	ORF 35 -36 -37 and	Del 36-37F2 (A, S)	ATACGATGCGTCGGTAGAGC	989, 384 or no	(Renault et al., 2012)
(605bpDel)	52968	38	Del 36-37R (A, S)	CGAGAACCCCATTCCTGTAA	amplification	
			C1590 (S)	CATGGTGATGAATGAAG		This paper
			C1591 (S)	CTTCATTCATCACCATG		This paper
			C1592 (S)	CATTCCTGTAAACAC		This paper
4	72414	Between ORF49 and	NC1 (A, S)	ACACCTAATGACCCCAAAGG	506	(Batista et al., 2015)
(NC1NC2 region)	72919	50	NC2 (A, S)	GACCAATCACCAGCTCAACA		
5	17402	ORF 11	ORF11For (A, S)	ACCACCGCGCCAAAATCTG	2116 or 731	(Martenot et al., 2013)
	19518		ORF11Rev (A, S)	CGCTTCCTATCACCTTGTGG		
			C1872 (S)	CTAGTCGTGCTCGTTCCTCTGC		This paper
			C1873 (S)	GGCAGAGATAGAACACAATG		This paper
			C1874 (S)	CATTGTGTTCTATCTCTGCC		This paper
6	133088	ORF 88a	ORF88aFor (A,S)	CCCAGTCTATTATCCAGGTAC	1020	(Martenot et al., 2013)
	134107		ORF88aRev (A,S)	ACCGTTCCTCAATCAGTCCC		

<sup>a</sup> Position on the OsHV-1 genome relative to the OsHV-1 reference genome, GenBank: AY509253

<sup>b</sup> Purpose: A amplification, S sequencing

at 72°C for 5 min. For the remaining regions, each reaction contained 10µl 5X HiFi reaction buffer (Bioline) (containing 10mM Mg2+), 0.5µl dNTP mix (100mM), 1µl each forward and reverse primer (400nM), 1µl (2U) Velocity DNA polymerase (Bioline), 1.5µl DMSO (BIO-21098 -Bioline Aust Pty. Ltd), 5µl template DNA and nuclease-free water to a final volume of 50µl; thermocycling was performed according to the following conditions: 1 cycle at 98°C for 2min; 35 cycles consisted in 98°C for 30s, 62°C for 30s and 72°C for 30s, and a final extension at 72°C for 10 min.

The amplicons were visualised using  $5\mu$ l of each amplification product on a 2% agarose gel, assessed against a molecular weight marker and visualised using RedSafe (iNtRON Biotechnology) on a GelDoc transilluminator (Biorad).

#### 2.3.3. Sequencing

Amplified PCR product was purified by incubation with ExoSAP-IT/ Cleansweep (ThermoFisher Scientific) then submitted for Sanger sequencing. For several samples where a single amplicon was not obtained, bands of the expected size were excised from the agarose gel and DNA was purified using the QIAquick Gel Extraction Kit (Qiagen). The amplicons were sent to the Australian Genome Research Facility or to the Monash Health Translation Precinct Medical Genomics Facility Australia for sequencing. Forward and reverse sequencing were performed in reactions with relevant PCR primers and internal primers (Table 2). Chromatograms were reviewed, analyzed and primer sequence was removed, using FinchTV (Geospiza).

#### 2.4. DNA data processing

For each region targeted, nucleotide sequences were aligned with the reference genome NC\_005881.2 and the two microvariant genomes KY242788 and KY271630, using MEGA10 (Kumar et al., 2018) with the MUSCLE algorithm. The sequences were trimmed and the chromatograms were again specifically checked for all mutations on forward and reverse strands. Base positions described in results are mapped to the reference genome NC\_005881.2.

Phylogenetic trees were constructed using MEGA 10 with the

Neighbour Joining method and 1000 replications of boostraps and the method based on Maximum Composite Likelihood was applied. Two trees were obtained: the first was based on the concatenated sequence of Regions 1, 2, 3, 4 and 6 (Region 5 was not included due to too few sequences - see results); the second tree was based on Region 1. The trees were unrooted because the ancestry of the reference genotype is uncertain. In addition, the use of another *malacoherpesviridae*, Haliotid herpesvirus, was impracticable because the ORFs had no sequence in common making alignment impossible. The unrooted tree allows the analysis of diversity but does not orient the genotypes temporally.

Nucleotide diversity (*Pi*) for Region 1 (C region) including the microsatellite region was calculated using DnaSP v6: DNA Sequence Polymorphism Analysis of Large Datasets (Rozas et al., 2017) and compared with the results of Mineur et al. (2015), after adding six sequences (JN800089, JN800075, JN800082, JN800083, JN800088 and JN800072) to be consistent with these authors. Briefly, the calculation was performed on the alignment; the indel polymorphism module with the "Multiallelic"gap option was used to consider all InDel events. Nucleotide diversity was listed according to geographic region. *Pi* values reported by Mineur et al. (2015) were corrected for frequency equal to one because multiple occurrences of an identical genotype may not be reported. To make the data comparable, *Pi* in the current study was not corrected because only one sequence per genotype was included in the calculation.

#### 2.5. Amino acid sequence prediction

In order to estimate whether some putative mutations could induce a modification in the phenotype of the variant, a prediction of amino acid sequence was performed on coding sites in Regions 1, 2, 3, 5 and 6. The translations of the ORFs were performed with ORF Finder (https://www.ncbi.nlm.nih.gov/orffinder/) and the Bioedit sequence alignment editor. The translations were checked using Blastp (https://blast.ncbi. nlm.nih.gov/Blast.cgi?PAGE=Proteins). These predictions were aligned with the prediction of the reference genome NC\_005881.2 and the two microvariant genomes KY242788 and KY271630 using Mega 10.

#### 3. Results

#### 3.1. DNA sequencing analysis

#### 3.1.1. Description of the variants

All nucleotide sequences are available on the NCBI database and accession numbers are listed in Table 3. Results and metadata for each of the 100 samples is provided in Supplementary data 1. Sequence alignments are illustrated in Supplementary data 2.

For Region 1, located at ORF 4, nucleotide sequence was obtained from 87 of 100 samples. None of the sequences were identical to the OsHV-1 reference sequence but they were partially similar to the microvariants, with the presence of most of the mutations except the two substitutions of adenine instead of guanine in the downstream region (bases 178,696 and 178,702). However, they all presented a novel substitution of an adenine instead of a guanine in a different position (base 178,705), also in the downstream region. Several mutations were observed allowing the identification of five different nucleotide sequences: G1, G2, G4, G6 and G7 (Fig. 1, Supplementary data 2). Eight of the sequences that grouped in G4 (Supplementary data 2) had 15 deletions (bases 178,558–178,572) in the microsatellite zone instead of the 12 deletions seen in microvariants (Burioli et al., 2017; Segarra et al., 2010), as was observed in OsHV-1 $\mu$ Var $\Delta$  15 reported in Martenot et al. (2012).

For Region 2, situated at ORF 43, nucleotide sequence was obtained from 77 of 100 samples. They were all different from the reference sequence and the microvariants. They were separated into two groups, classified by one deletion of a thymine in the downstream region (base 60,504) (G8) for 36 samples and two deletions of two adenine in the upstream region (bases 60,059 - 60,060) for 41 samples (G9) (Fig. 2, Supplementary data 2).

In the third region coding for ORF 35 to ORF 38, nucleotide sequences were obtained from 96 of 100 samples and all were identical to the two microvariant sequences. The features included the large deletion zone of 606bp (bases 52,251–52,856) common to microvariants and the

#### Table 3

GenBank accession nun	bers of the 29 DNA	groups of t	he 6 regions analysed.
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					• •
Region		ion	Gene	Sequence	GenBank
	0			Group	Accession No.
	1	C region	ORF4	G1	ON953753
		-		G2	ON953754
				G4	ON953755
				G6	ON953752
				G7	ON953756
	2	A region	ORF 42-43	G8	ON953757
				G9	ON953758
	3	605 bpDel	ORF 35 to 38	G10	ON953759
	4	NC2	Non-coding, between ORF	G11	ON954001
		region	49 and ORF 50	G12	ON954014
				G13	ON954007
				G14	ON954004
				G15	ON954003
				G17	ON954002
				G18	ON954013
				G19	ON954012
				G20	ON954006
				G21	ON954010
				G22	ON954009
				G23	ON954008
				G24	ON954005
				G29	ON954011
	5	ORF 11	ORF 11	G31	ON953762
		gene		G32	ON953760
				G33	ON953763
				G34	ON953761
	6		ORF 88	G35	ON953765
				G36	ON953764
				C37	ON053766

substitution of a cytosine instead of a thymine (base 52,885). They were classified into G10 (Fig. 3, Supplementary data 2).

Eighty-seven of 100 samples presented an amplification of the fourth region located on the non-coding site between ORF 49 and ORF 50. All the nucleotide sequences presented a common substitution of an adenine instead of a guanine in the upstream zone (base 72,489) and two deletions of two thymine next to the middle zone (bases 75,592–75,593). However there were many differences compared to the reference genotype and microvariants, consisting of 19 mutations over 408 nucleotides which were not observed in the microvariant genotypes. These mutations allowed the characterization of 14 groups (Fig. 4, Supplementary data 2), confirming the prior observations of high variability in this region (Batista et al., 2015). Four samples of a nucleotide sequence named G12 presented a thymine as well as a cytosine (Y) at the same position (base 72,536). The chromatograms of this and all the other regions of these four samples comprising G12 were unambiguous. All samples which contained the G12 variant were pools of 5 spat (Supplementary data 1).

For Region 5 sitting on ORF 11, nucleotide sequence was obtained from 42 of 100 samples. None had the large deletion observed in the microvariants. Even though the nucleotide sequences were close to the reference sequence, a few mutations were present allowing their classification into four groups (G31, G32, G33, G34) (Fig. 5, Supplementary data 2). All had a common substitution of a thymine instead of a guanine (base 18,621).

Of the 91 samples which amplified from Region 6 in the first part of ORF 88, 54 nucleotide sequences were similar to the reference sequence (G37). The others clustered into two groups, comprising the substitution of a cytosine instead of a guanine (base 133,574) in 36 samples (G36) and a substitution of a guanine instead of an adenine (base 133,574) in one sample (G35) (Fig. 6, Supplementary data 2).

Nucleotide sequences for Regions 1, 2, 3, 4 and 6 were obtained from 61 samples, which when concatenated led to the identification of 12 DNA variants (Table 4). V1 and V8 were the most numerous of these variants.

#### 3.1.2. Phylogenetic analyses

Firstly, a phylogenetic tree using Regions 1, 2, 3, 4 and 6 was constructed with the concatenated nucleotide sequences of the 12 variants and the 29 whole genomes present in NCBI. All Australian genotypes clustered in the same clade, distinct from isolates from Europe, China and the reference genotype and with a distinctly shorter branch length (Fig. 2). Secondly, a phylogenetic tree was constructed based only on the C region (Region 1), which has been well-studied internationally, in order to incorporate a wider range of isolates (Fig. 3). A single cluster distinguished all Australian genotypes from those from Europe, New

Table 4

. Description of the 12 genotypic variants based on concatenated nucleotide sequences of Regions 1,2,3,4 and 6 for 61 isolates according to their group identification numbers for these regions.

				0		
Variant	n	R1 ORF 4	R2 ORF 43	R3 ORFs 35, 36, 37, 38	R4 uncoded site between ORF49 and 50	R6 ORF 88
V1	32	G7	G8	G10	G11	G36
V2	3	G1	G9	G10	G11	G37
V3	1	G7	G9	G10	G20	G37
V4	1	G7	G9	G10	G21	G37
V5	1	G7	G9	G10	G29	G37
V6	2	G7	G8	G10	G11	G37
V7	5	G7	G9	G10	G13	G37
V8	10	G7	G9	G10	G11	G37
V9	2	G4	G9	G10	G12	G37
V10	2	G7	G9	G10	G14	G37
V11	1	G7	G9	G10	G15	G37
V12	1	G7	G9	G10	G17	G37



Fig. 2. Distribution of the DNA variants according to the NCBI deposited nucleotide sequences. The unrooted phylogenetic tree was performed using the neighbour joining method and 1000 bootstrap replications were applied based on Method based on Maximum Composit Likehood. Regions 1,2,3,4 and 6 were concatenated on 61 Australian isolates showing 12 variants (V1, V2, V3, V4, V5, V6, V7, V8, V9, V10, V11, V12). Yellow rectangle is the reference nucleotide sequence of OsHV-1. Metadata for the nucleotide sequences of the current study that are included in the tree are in Supplementary data 1 while metadata for the nucleotide sequences that were already available in NCBI are in Table 1, Supplementary data 3.

Zealand, Japan and the Americas.

#### 3.1.3. Diversity

The number of genotypes available for analysis had almost doubled from 48 to 95 since the study of Mineur et al in 2015 (Table 5). The Australian isolates had the lowest genetic diversity overall (*Pi* 0.99 ×  $10^{-3}$ ) when compared to viruses from other regions of the world where diversity ranged from  $5.31 \times 10^{-3}$  to  $7.61 \times 10^{-3}$  (Table 5).

#### 3.1.3. Influencing factors

The distribution among the sites over time of the 12 DNA variants

identified from concatenated nucleotide sequences is illustrated in Table 6 while similar data for all genomic regions is presented in Table 1, Supplementary data 2.

Considering the concatenated nucleotide sequences, up to four variants were found in one summer season in the Georges River, compared to only one or two variants in the Hawkesbury River. Nine of the 12 variants occurred at only one site and in only one summer season. The variants V1 and V6 were present only in the Hawkesbury River; V1 was present at most sites and in each of the four summers, while V6 was found only at Kimmerikong Bay and Mooney Mooney in summer 2013–2014. The other variants were confined to the Georges River



Fig. 3. Distribution of the DNA variants according to nucleotide sequences deposited in NCBI. The unrooted phylogenetic tree was performed using the neighbour joining method and 1000 bootstrap replications were applied based on maximum composite likelihood. Region 1 from 87 Australian isolates was used, showing 5 groups (G1, G2, G4, G6, G7). Crosses indicate Australian nucleotide sequences extant in NCBI; asterisks indicate the New Zealand nucleotide sequences. Yellow rectangle is the reference nucleotide sequence of OsHV-1. Metadata for the nucleotide sequences of the current study that are included in the tree are in Supplementary data 1 while metadata for the nucleotide sequences that were already available in NCBI are in Table 2, Supplementary data 3.

where V8 was identified at various sites within Woolooware Bay in three summer seasons. The other variants in the Georges River were detected only at a single site and a single sampling date.

More than one variant was detected over time at some sites; for example, five were seen at Site B in the Georges River while two were seen at both Kimmerikong Bay and Mooney Mooney in the Hawkesbury river (Table 7). More than one variant was found in the same summer season at some sites; for example, in 2014–2015 V8, V11 and V12 were found at Site B, while in 2013-2014 both V1 and V6 were found at

Kimmerikong Bay and at Mooney Mooney. Furthermore, there were also several instances where different variants were detected in the same population of oysters at the same time: V11 and V12 were both present at Site B on 27/1/15; V1 and V6 were both present at Kimmerong Bay and Mooney Mooney on 13/12/13 and also on 25/1/14.

There were similar observations for Region 4 (Fig. 4, Supplementary data 2), which had the most variability; 12 of the 14 variants were found in only one season and at only one site. Three of five variants in Region 1 and two of four variants in Region 5 were also seen in only one season

Nucleotide diversity in the C region (including the microsatellite region) of OsHV-1 from different geographical regions. The data are based on sequences listed in Table 2 of Supplementary data 3 as well as sequences from the current study. Six sequences were added to be consistent with Mineur et al (2015) (see materials and methods). NW Pacific includes isolates from China, Japan and Korea; NE Pacific includes Mexico and USA; Europe includes France, Ireland, Italy, Portugal and Spain; Australasia includes Australia and New Zealand.

	Current stud	у	Mineur et al (2015)			
Geographic region	No. of genotypes	Nucleotide diversity ( <i>Pi</i> )	No. of genotypes	Nucleotide diversity ( <i>Pi</i> ) **		
All	95	$7.46 imes10^{-3}$	48	$14.97\times10^{-3}$		
NW Pacific	45	$6.45  imes 10^{-3}$	27	$9.34 imes10^{-3}$		
Australasia	10	$1.49  imes 10^{-3}$	3	$2.09 imes10^{-3}$		
Australia	7	$0.99 imes10^{-3}$	nr	nr		
NE Pacific	5	$6.24 imes10^{-3}$	2	$6.27 imes10^{-3}$		
Europe all vears	41	$\textbf{7.61}\times 10^{-3}$	16	$15.30\times10^{-3}$		
*Europe < 2008	9	$\textbf{3.43}\times 10^{-3}$	10	$\textbf{4.67}\times 10^{-3}$		
Europe $\geq$ 2008	32	$2.07\times10^{-3}$	6	$3.45\times10^{-3}$		
France	27	$5.31\times10^{-3}$	nr	nr		

\*One sequence (AY459363) with a high frequency of undetermined nucleotides ("N") was excluded from the current study

\*\*corrected for Hd=1, see materials and methods

nr, no result

#### and at only one site.

Two isolates from Tasmania were not incorporated into the phylogenetic analysis because of incomplete sequencing of regions from these samples. However one of these samples was the only isolate classified as G6 in Region 1 (Table 1 and Fig. 1, Supplementary data 2); there were no sequencing data from this region of the other sample from Tasmania. The oysters sampled in New South Wales had been supplied from at least nine different production batches (i.e. different broodstock spawnings) from one hatchery in Tasmania. Oysters grown from five of these batches yielded more than one variant, and in one case, five different OsHV-1 variants, which has important implications when assessing the sources of the virus (see discussion).

#### 3.2. Amino acid sequence prediction of variants

#### 3.2.1. Description of the predicted amino acid sequences

Predicted amino acid sequences were aligned with the reference and microvariant sequences. All alignments are illustrated in Supplementary data 2. Regions 3 and 4 were not included in this analysis because Region 3 had a large deletion of ORF 36 and 37, inducing overlap of ORF 35 and 38 and creating uncertainty about whether a longer ORF could be translated, while Region 4 was a non-coding area.

For Region 1, two amino acid sequences were identified for ORF 4 among the five DNA sequences described above in part 3.1.1. They were both different to the reference genotype and the microvariants. The first group was named <<A>> and contained G1, G2, G4 and G7; it had an asparagine (N) instead of an aspartic acid (D). The second group named <<B>> contained G6 and had in addition a leucine (L) instead of a serine (S) (Fig. 7, Supplementary data 2).

For Region 2, the two DNA groups characterised for ORF 43 presented two different amino acid sequences. The first was named <<C>> and contained G8. It had a similar amino acid sequence to the reference genotype and the microvariant. The second was named <C>>. It contained G9 and was characterised by three successive mutations of a serine (S), lysine (K) and a deletion instead of a phenylalanine (F), a glutamic acid (E) and an arginine (R) (Fig. 8, Supplementary data 2).

Region 5 located on ORF 11 had four DNA groups which translated into 3 amino acid sequences. One was similar to OsHV-1 reference genotype, and was named « E ». The second was named « F » and had a mutation of an asparagine (N) instead of a lysine (K). The third was named « G » and had 4 substitutions of a lysine (K) (Fig. 9, Supplementary data 2).

Region 6 coding for the first part of ORF 88 presented two amino acid sequences named « H » containing G36 and G37 and « I » containing G35. <<H>> had a similar sequence to the reference genotype and one of the microvariants, KY242785, while <<I>> had one mutation of a glutamic acid (E) instead of lysine (K) (Fig. 10, Supplementary data 2).

The amino acid prediction of the 12 DNA variants identified among 61 samples resulted in two profiles based on ORF 4, ORF 43 and ORF88: ACH (n = 34) and ADH (n = 27) (Table 8). Region 5 (ORF11) was not included because there were only 42 sequences available for it.

#### 3.2.2. Influencing factors

The amino acid profile of isolates was different between the two estuaries in NSW (Tables 6 and 7). The ACH profile was found only in the Hawkesbury River, an estuary located 40 km north of Sydney while the ADH profile was found only in the Georges River, an estuary located 20 km south of Sydney.

#### 4. Discussion

## 4.1. Australian OsHV-1 isolates have lower diversity than OsHV-1 isolates globally

It is difficult to compare genomic diversity between published studies of OsHV-1 due to differences in the genomic regions sequenced, sample sizes, time span, geographic extent, host species and disease state (healthy, sick, or dead). However, in this study we report diversity of *Ostreid herpes virus*1 in estuaries near Sydney Australia that may be lower than in other geographic regions of the world.

Regions of the OsHV-1 genome under selection pressure and other regions which accumulate a high number of substitutions have been reported (Morga et al., 2021). Selection pressure is less on a non-coding area and mutations acquired over time are less likely to be eliminated. The diversity revealed in this study is mainly explained by inclusion of non-coding Region 4 between ORF 49 and ORF 50, which is a polymorphic area (Batista et al., 2015). In this region alone, 14 groups were identified among the 87 samples (one variant for every 6.2 isolates). Similarly, Batista et al. (2015) identified one variant for every 6.0 isolates examined in a study of 18 isolates from mostly diseased *C. gigas* in Portugal.

C region, corresponding to Region 1 in the present study, is the most commonly analysed part of the OsHV-1 genome. With the typical primers C2/C6 used for amplification there is representation of part of a non-coding region and part of ORF 4 (Batista et al., 2015). Nucleotide diversity in the C region was calculated formally and compared to data published in 2015 (Mineur et al., 2015). There were more sequences in public databases, but the nucleotide diversity had not changed markedly except for Europe where it appeared to have decreased (from 15.3  $\times$  $10^{-3}$  to 7.6  $\times$   $10^{-3}$ ). The Australian isolates had the lowest genetic diversity (*Pi*  $0.99 \times 10^{-3}$ ) overall. Globally, 48 different genotypes had been identified in C region (including the microsatellite region) by 2015 (Mineur et al., 2015). Expressed in simple terms of the frequency of genotypes among isolates examined, in Australia seven genotypes have been identified among 89 samples in diseased C. gigas (five from 87 isolates in the present study and two existing in NCBI), which equates to one variant for every 12.7 isolates. In contrast, in a study covering a wide area of coastal Japan, 23 genotypes were found among 123 samples from healthy Crassostrea species (one variant for every 5.4 isolates) (Shimahara et al., 2012); in both countries samples were collected over about 5 years. The genetic diversity among diseased oysters in Australia was clearly much less than that in healthy oysters in Japan. The data from Japan in the north west Pacific region likely represent the diverse environmental reservoir of viral types detectable in healthy shellfish,

Distribution of 12 genomic variants of OsHV-1 among 61 isolates in the Georges and Hawkesbury River estuaries according to site and date of collection over five summer seasons between 2011-2012 and 2015–2016.

	2011-20	)12			2012-2	013			2013-20	014			2014-2	015			2015-2	016		
	Isolate	Site	Date collected	Variant	Isolate	Site	Date collected	Variant	Isolate	Site	Date collected	Variant	Isolate	Site	Date collected	Variant	Isolate	Site	Date collected	Variant
Georges River																				
0	13	Site C	16-11-11	V9	19	Site A	07-11-12	V4	65	The Shed	28-01-14	V7	102	Neverfail	16-12-14	V10	108	Site C	09-11-15	V8
	14	Site C	16-11-11	V9	26	Site B	28-11-12	V5	66	The Shed	28-01-14	V7	103	Neverfail	16-12-14	V10	109	Site C	09-11-15	V8
	4	Site A	24-11-11	V8					67	The Shed	28-01-14	V7	104	Site B	27-1-15	V11				
	9	Site B	10-02-12	V3					68	The Shed	28-01-14	V7	105	Site B	27-1-15	V12				
									69	The Shed	28-01-14	V7	79	Site C	27-1-15	V8				
									70	Site C	25-02-14	V2	80	Site C	27-1-15	V8				
									71	Site C	25-02-14	V2	81	Site C	27-1-15	V8				
									73	Site C	25-02-14	V2	82	Site C	27-1-15	V8				
													84	Site C	27-1-15	V8				
													85	Site C	27-1-15	V8				
														106	Site B	24-2-	V8			
																15				
Hawkesbury River					31	Mullet	21-01-13	V1	42	Kimmerikong	31-12-13	V1	74	Mooney	18-11-14	V1	114	Mullet	26-10-15	V1
					32	Mullet	21-01-13	V1	43	Kimmerikong	31-12-13	V1	75	Mooney	18-11-14	V1				
					33	Mullet	21-01-13	V1	44	Kimmerikong	31-12-13	V1	76	Mooney	18-11-14	V1				
					34	Mullet	21-01-13	V1	46	Kimmerikong	31-12-13	V6	77	Mooney	18-11-14	V1				
					36	Mullet	21-01-13	V1	48	Mooney	25-01-14	V1	78	Mooney	18-11-14	V1				
					37	Mullet	21-01-13	V1	49	Mooney	25-01-14	V1	110	Porto	28-11-14	V1				
					38	Mullet	21-01-13	V1	50	Mooney	25-01-14	V1	111	Porto	28-11-14	V1				
					41	Coba	29-01-13	V1	51	Mooney	25-01-14	V1	112	Porto	28-11-14	V1				
					39	Porto	15-02-13	V1	52	Mooney	25-01-14	V1	113	Porto	28-11-14	V1				
					40	Porto	15-02-13	V1	53	Mooney	25-01-14	V1								
									54	Mooney	25-01-14	V1								
									55	Mooney	25-01-14	V1								
									56	Mooney	25-01-14	V1								
									57	Mooney	25-01-14	V6								

Distribution of 12 DNA variants and protein prediction profiles among 61 samples from ten different sites in the Georges and Hawkesbury river estuaries between 2011 and 2016.

Estuary	Site	No. samples	DNA variant	Predicted protein
Hawkesbury	Coba Bay	1	V1	ACH
river ( $n = 34$ )	Kimmerikong	4	V1 ( <i>n</i> = 3) V6	ACH
	Bay		(n = 1)	
	Mooney	15	V1 ( <i>n</i> = 14) V6	ACH
	Mooney		(n = 1)	
	Mullet Creek	8	V1	ACH
	Porto Bay	6	V1	ACH
George river (n	Neverfail	2	V10	ADH
= 27)	Site A	2	V4 V8	ADH
	Site B	5	V3 V5 V8 V11	ADH
			V12	
	Site C	13	V8 (n = 8) V9	ADH
			(n = 2) V2 $(n$	
			= 3)	
	The shed	5	V7	ADH

#### Table 8

Protein profiles of the 12 DNA variants identified among 61 samples from ten different sites in the Georges and Hawkesbury river estuaries between 2011 and 2016.

Variant	n	ORF 4	ORF 43	ORF 88
V1	32	А	С	Н
V2	3	Α	D	Н
V3	1	Α	D	Н
V4	1	Α	D	Н
V5	1	Α	D	Н
V6	2	Α	С	Н
V7	5	Α	D	Н
V8	10	Α	D	Н
V9	2	Α	D	Н
V10	2	Α	D	Н
V11	1	Α	D	Н
V12	1	Α	D	Н

while the low diversity in Australia may represent selected environmental strains of OsHV-1 capable of causing disease in farmed oysters under certain environmental conditions, that then became locally dominant. Relatively low diversity was seen in Europe after 2008 when OsHV-1  $\mu$ *Var* became the most common strain in diseased oysters (Table 5). In Europe there had been spread of the virus with movements of live oysters for farming, for example from France to Ireland and Jersey (Peeler et al., 2012). Within France, OsHV-1  $\mu$ *Var* was dispersed widely with commercial movements of live oysters from Marennes-Oléron to farming areas on both the Atlantic and Mediterranean coasts (Delmotte et al., 2022). Thus, in addition to the health/disease condition of oysters and their geographic locations, the observed regional differences in the diversity of OsHV-1 genotypes depends on epidemiological differences between the sources and modes of spread of the viruses. All these factors will need to be considered in future studies of the diversity of OsHV-1.

In Region 4 we observed a double nucleotide (thymine and cytosine) in four individual samples for which remaining nucleotide sequences were unambiguous (G12 – Fig. 4, and Supplementary data 2) and otherwise matched the nucleotide sequence of G18 which had a cytosine at this position (Fig. 4, Supplementary data 2). Our first hypothesis is that there has been a recent mutation at this position to explain why we found two different copies of this part of the genome, one being the ancestral form and the other being the mutated one. This is feasible due to the persistence of the herpesvirus in surviving oyster hosts (Evans et al., 2017b), because gene exchange in herpesviruses can occur by recombination (Thiry et al., 2005) and because Rosani and Venier (2017) showed the presence of DNA recombination-initiating promoter binding in *Malacoherpesviridae*. The four samples with this mutation

were from oysters in close proximity to one another and they could have been infected almost simultaneously from a host in which the mutation first occured, resulting in a pool of two variants in each of the four samples. This is consistent with Martenot et al (2011) who reported various genotypes in the same batch of oysters and Morga et al (2021) who reported various genotypes in the same individual. A second possibility would be that there were two otherwise identical but long-standing OsHV-1 variants in each of the four samples, due to different individuals containing a different genotype of OsHV-1, because each sample came from a pool of five individual oysters. While this does increase the chance of mixing several variants into one DNA extract, in no other sample did we find a group with a thymine at this position. The least likely explanation because the non-coding area between ORF 49 and ORF 50 is present as a single copy in the complete OsHV-1 genomes available on NCBI, is the existence of a double copy of this region.

The microsatellite zone of the C region of OsHV-1 is polymorphic because of repeated trinucleotides which allow replication mistakes, as observed by many authors (Martenot et al., 2012; Martenot et al., 2011; Mineur et al., 2015; Renault et al., 2014). The range in the number of ACT trinucleotide repeats across all genotypes globally was reported to be 3 - 13 (Mineur et al., 2015), corresponding to a range of 3–10 trinucleotide deletions (9–30 base deletions). However, we observed only two profiles among a large number of isolates, either 12 or 15 base deletions. This is consistent with the only other Australian sequences deposited in NCBI; these came from the first episode of mortality in 2010 and had 12 base deletions.

#### 4.2. Australian isolates differed from OsHV-1 isolates globally

Concatenated nucleotide sequences showed a distinct cluster including all the Australian isolates, different from all other genotypes reported worldwide and supported by the distinct distribution with a shorter branch size for the Australian genotypes (Fig. 2). Although the bootstrap value was only 63, this could be explained by the presence of Region 3 located between ORF 35 and 38 including a large deletion area and region 5 located on ORF 11, with a 462bp deletion only in the microvariant. These differences are calculated as multiple mutation events when it is more likely that each corresponds to a single mutation event. Burioli et al. (2016) studied the same disturbed region and discussed the need for sophisticated algorithms for alignment and phylogenetical analysis, adapted to take into account DNA-virus evolution analysis, in order to obtain accurate results with the correct phylogenetic distance. The Australian clade was also confirmed by the inclusion of the only two NCBI sequences from Australia, KC685525.1 and MT157286.1 (Burge et al., 2020; Jenkins et al., 2013) in the C region tree (Fig. 3). This is consistent with the phylogeny reported in Mineur et al. (2015), which was performed on the same locus with another algorithm, median joining genotype, and showed three closely related genotypes from temperate Australasia. In contrast, the phylogenetic tree presented in Burge et al. (2021) with the neigbour joining method showed that the Australian genotype was close to the Japanese, American, Korean and French microvariants. However, unlike the current phylogeny, the tree described in Burge et al. (2021) is not based on evolutionary distance and contains too few samples - a single genotype from Australia - which is not sufficient to construct a cluster. The Australian viruses appear to be genetically different across the genomic regions analyzed and this raises the question about their true phylogenetic origins. Sanger sequencing of a limited number of genomic regions may bias analyses compared to whole genome analysis, which will be indispensable for determining clades of OsHV-1 and should be the subject of future studies.

## 4.3. Australian isolates were not OsHV-1 reference strain, OsHV-1 $\mu$ Var or microvariants

Neither OsHV-1 reference genotype nor OsHV-1 µVar were identified

among the Australian samples. Currently OsHV-1 "microvariants" are described as being close to OsHV-1 µVar as defined by Segarra et al. (2010) but not identical to it (Burge et al., 2021; Morga et al., 2021). In all of the Australian isolates the two regions which characterise OsHV-1 µVar according to Segarra et al. (2010), ORF 4 and ORF 43, had dissimilar nucleotide sequences. None of these sequences had the large deletion in ORF 11 reported in the microvariant sequences (Burioli et al., 2017; Martenot et al., 2013; Morga et al., 2021) and presented a similar nucleotide sequence to the OsHV-1 reference virus on ORF 11 except for one point mutation. Moreover, half the samples from Australia did not present an amplification product from this region. Since there was enough biological material in the same samples to perform sequencing of the other regions, we suggest that there were changes at the primer binding sites, which others have also mentioned to be a likely reason for this problem (Arzul et al., 2001b; Martenot et al., 2011); PCR-sequencing of the other regions was not uniformly successful either, consistent with many other studies of OsHV-1.

The variant of OsHV-1 known as  $\mu$ Var is thought to be responsible for the sudden escalation of mass mortality events in *C. gigas* in France from 2008, and then later in other European countries. However,  $\mu$ Var and the more recently described "microvariants" are clearly not the only variants of OsHV-1 associated with mass mortality events in *C. gigas*. The disease outbreaks in Australia since 2010 have not been caused by these types of OsHV-1 but have been just as sudden in onset, recurrent and economically devastating for the oyster industry as those in France (Fuhrmann et al., 2019). Therefore further research is required to understand the virulence determinants of OsHV-1 and how these are coded in the genome.

#### 4.4. Internal comparison of Australian isolates

The two estuaries studied in New South Wales had different OsHV-1 profiles. At the Georges River sites, ten variants were identified over five summer seasons while at the Hawkesbury River sites only two variants were found. Delmotte et al. (2022) showed a great diversity of variants in areas with high oyster farming activity. However, the causes of the different patterns of distribution of variants here are unknown since these two estuaries in Australia had equivalent, intense oyster farming activity using similar genetic stock of C. gigas from the same OsHV-1 free hatchery in Tasmania. As a consequence of the two distributions of DNA variants, the amino acid sequences of the viruses present in the Hawkesbury and Georges river estuaries were different: the ACH profile in the Hawkesbury River and ADH in the Georges River. This distinct pattern has a precedent in Europe, but with far greater geographic separation than the 60 km that exists between these two estuaries. Based on RNA-Seq SNP analysis, Delmotte et al. (2020) reported two genetically different OsHV-1 populations in POMS outbreaks in C. gigas on the Atlantic and Mediterranean coasts of France; they had amino acid sequence differences in viral membrane proteins, which may affect virulence (Delmotte et al., 2020).

Due to the selection of samples based on high viral load, which in general is correlated with mortality (Oden et al., 2011), it was not appropriate to compare the OsHV-1 isolates in this study according to their degree of replication fitness or observed mortalities. However, in the oyster populations from which the samples were collected there were differences in the patterns of mortality: spat in the Georges River estuary were 6.4 times more likely to have mortality due to OsHV-1 than those in the Hawkesbury River estuary (Whittington et al., 2019). Subclinical infection of oysters with OsHV-1 was also more likely in the Georges River (Evans et al., 2017b; Whittington et al., 2019). It is possible that the mutually exclusive genotypic variants of OsHV-1 identified between the two estuaries differed in virulence. However, other factors could also be involved. While oyster genotype was similar between the two rivers because of the common origin of spat, there could be a wide range of environmental differences. These could include differences in the co-occurrence of reservoirs/hosts, paticulate vectors,

filter feeding species and seaweeds that could all influence the severity of POMS (Dugeny et al., 2022; Evans et al., 2017c; Pernet et al., 2021; Whittington et al., 2018).

To our knowledge there were no transfers of commercial oysters between the Georges and Hawkesbury Rivers after first detection of OsHV-1 in the Georges River late in 2010 due to government biosecurity directives (Paul-Pont et al., 2014). The only common factor between the two oyster production locations was the recruitment site of all batches of farmed C. gigas in this study: Tasmania. However, the current data indicate that the virus contamination did not come from this recruitment step but from the culture environment. There are three reasons for this: (i) consistent pre-shipment PCR testing of all batches of spat from the hatchery by the government laboratory in Tasmania and again upon arival in NSW by the researchers (all negative results) means that spat were specific-pathogen free (SPF); (ii) different genotypes of OsHV-1 were detected in the same batch of spat after deployment in NSW; (iii) a unique genotype (G6) of OsHV-1 was detected in Tasmania; G6 was not found among 85 samples from the Hawkesbury River or Georges River. Epidemiological observations alone had suggested that recurrent POMS outbreaks in Australia were derived from the environment and were not due to disease transfer with ovsters during farming (Whittington et al., 2018), a feature which is now strongly supported by DNA sequence evidence. This scenario is in direct contrast to France where evidence points to a radiation of disease associated with commercial spat distribution (Delmotte et al., 2022). The time of first appearance of OsHV-1 in a putative environmental host in Australia cannot be determined, but it could have been a point source introduction to a naive population, then adaptation to local conditions. In fact there are historical records of a virus like Ostreid herpesvirus from Ostrea angasi in Western Australia (Hine and Thorne, 1997). These authors demonstrated herpes-like viral particles by electron microscopy, but did not have the molecular tools to identify the virus. In any case, this observation raises the possibility that OsHV-1 could be either indigenous to Australia or could have been introduced many decades ago and become endemic, and it could have mutated to form its own lineage over time, as is supported by the current phylogenetic analysis. Knowing that OsHV-1  $\mu Var$  emerged around 2008 in Europe and then dispersed rapidly, the absence of OsHV-1  $\mu$ Var in Australia suggests that there has been effective compliance with the strict quarantine laws that aim to prevent introduction of exotic pathogens into Australia.

Arzul et al. (2001a) speculated 20 years ago that pathogenic forms of OsHV-1 arose from possibly benign bivalve herpesviruses which could be transmitted between bivalve species and which would be favoured by modern shellfish farming practices (Arzul et al., 2001a; Arzul et al., 2001b). In Australia, there is evidence that invertebrate species in the Georges River besides C. gigas are exposed to OsHV-1 and may become infected, but their role in the epidemiology of POMS remains to be determined (Evans et al., 2017c). The environment is an important factor in the transmission cycle because OsHV-1 can be found in seawater and sediments (Evans et al., 2014; 2017) and is likely transported on suspended particles over 5µm (Evans et al., 2017a; Evans et al., 2017c; Paul-Pont et al., 2013b; Whittington et al., 2015b). Depending on conditions such as temperature, the virus is expected to persist in seawater and remain infectious for several days according to in vitro experimentation (Hick et al., 2016; Martenot et al., 2015). This gives the virus time to transit between hosts through the environment. Ostreid herpesvirus 1 can infect several different host species (Arzul et al., 2001a; Bai et al., 2016; Batista et al., 2015; Bookelaar et al., 2018; Comps and Cochennec, 1993; Evans et al., 2017c; Farley et al., 1972; Hine and Thorne, 1997; Hine et al., 1998; Prado-Alvarez et al., 2021; Renault and Arzul, 2001; Renault et al., 2001; Xia et al., 2015) and may persist in survivors of outbreaks or remain latent (Arzul et al., 2002; Evans et al., 2017b), which may explain why each environment has its own assemblage of viruses. While the molecular evidence strongly suggested that the virus contamination came from reservoirs in the local culture environment rather than a point source such as an oyster

hatchery, the distribution of the variants across sites and time suggests that some reservoirs were persistent and widespread, while others were highly site and time specific and therefore possibly short-lived, or that transmission was infrequent. The reservoirs may be subject to local environmental influences which affect their distribution and abundance. Further field research is required to discover the environmental reservoir of OsHV-1 and the specific environmental factors that facilitate emergence of the virus. Climate change, short term climatic and water quality variations and intensification of farming are among the many factors to be considered.

#### 5. Conclusion

Sequencing analyses of OsHV-1 isolates from estuaries near Sydney, Australia, revealed a low diversity of genotypes relative to other geographic regions, reflecting differences in sources and modes of spread of the viruses and the types of oyster populations (healthy, or diseased) that have been examined so far. Importantly, Australian POMS outbreaks were associated with a type of OsHV-1 distinct from others globally and were not a "microvariant" genotype. Elsewhere, since 2008 OsHV-1 microvariants have been the only variants associated with mass mortality events in C. gigas. Future whole genome sequencing investigations will allow more detailed classification of Australian isolates of OsHV-1. Based on both DNA and amino acid sequences, OsHV-1 variants were absolutely correlated to estuary. The C. gigas populations in the two estuaries near Sydney had different prevalence and severity of POMS, which suggests that the geographically mutally exclusive genotypes differed in virulence. The molecular evidence supports field epidemiological evidence that the disease-causing virus came from reservoirs in the local environment. The sequencing findings in this study help inform understanding of the various sources and modes of spread of OsHV-1 that have led to mass mortality events in farmed C. gigas globally.

#### List of supplemental data

Supplemental data 1

Table 1 Metadata for all samples

Table 2 Citations of scientific publications describing the populations of oysters from which samples were collected

Supplemental data 2

Table 1 Variants by genomic region and geographic location Figs. 1 to 10 DNA and amino acid sequence alignments

Supplemental data 3

Genbank accession numbers of isolates included in phylogenetic trees

#### CRediT authorship contribution statement

Suzanne Trancart: Investigation, Data curation, Methodology, Visualization, Formal analysis, Writing – original draft, Writing – review & editing. Alison Tweedie: Investigation, Data curation, Writing – review & editing. Olivia Liu: Investigation, Writing – review & editing. Ika Paul-Pont: Conceptualization, Investigation, Writing – review & editing. Paul Hick: Investigation, Writing – review & editing. Maryline Houssin: Resources, Investigation, Supervision, Writing – review & editing. Richard J. Whittington: Funding acquisition, Project administration, Supervision, Resources, Conceptualization, Investigation, Data curation, Formal analysis, Writing – original draft, Writing – review & editing.

#### **Declaration of Competing Interest**

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

#### **Data Availability**

Comprehensive metadata for each sample are provided in Supplementary Data, linked to prior publications that contain more details. All sequences are available in GenBank.

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#### Supplementary materials

Supplementary material associated with this article can be found, in the online version, at doi:10.1016/j.virusres.2022.198994.

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