Low susceptibility of sockeye salmon *Oncorhynchus nerka* to viral hemorrhagic septicemia virus genotype IVa

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ABSTRACT: Viral hemorrhagic septicemia virus (VHSV) genotype IVa is an endemic pathogen to the marine waters of British Columbia, with numerous marine fishes being susceptible to infection and disease, including Atlantic salmon *Salmo salar* reared in open net-pen aquaculture. The susceptibility of Atlantic salmon and sockeye salmon *Oncorhynchus nerka* to VHSV-IVa infection was evaluated using exposure routes including injection, static immersion, and cohabitation with diseased Pacific herring *Clupea pallasii*. Exposed fish were monitored for mortality and external pathology, mortalities were tested by virus isolation assay, and live fish were regularly sampled and screened for infection. Among injected sockeye, VHSV was detected in 1 mortality (n = 195) and 2 sub-sampled fish (n = 30), whereas sockeye exposed by immersion and cohabitation did not experience mortality nor was systemic infection indicated by tissue screening. Injection and cohabitation exposure routes confirmed the susceptibility of Atlantic salmon to VHSV. Neither sockeye nor Atlantic salmon surviving the cohabitation served as a reservoir of VHSV, but Pacific herring did. The results suggest that VHSV-IVa poses low risk to sockeye salmon under natural routes of exposure.

KEY WORDS: Viral hemorrhagic septicemia virus · VHSV · Sockeye salmon · Susceptibility

1. INTRODUCTION

Sockeye salmon *Oncorhynchus nerka* is a keystone species in the North Pacific Ocean that is highly prized as a food commodity. Of the 7 species of Pacific salmon, sockeye salmon are the third most abundant after pink salmon *O. gorbuscha* and chum salmon *O. keta*. Due to their preferential use of lake habitats in juvenile rearing stages, the distribution of sockeye salmon is largely restricted to north temperate rivers with accessible lakes.

The Fraser River drainage of British Columbia, possessing extensive lake rearing areas, supports one of the largest spawning complexes of sockeye salmon in the North Pacific Rim (Burgner 1991). However, between the early 1990s and 2009, fisheries managers and biologists recognized a steady and profound decline in Fraser River sockeye productivity. In 2009, record low returns of sockeye salmon to the Fraser River forced closure of the fishery for the third consecutive year and prompted a federal investigation, the Commission of Inquiry into the Decline of Sockeye Salmon in the Fraser River. After review of numerous technical reports and testimonies of expert witnesses, the Commission concluded that sockeye salmon experience multiple stressors that may affect their health and their habitats at various stages of their life and suggested that multiple factors likely contributed to the decline (Cohen 2012). Moreover, the Commission highlighted knowledge gaps, uncer-
tainties, and future research needs to provide better information about the challenges faced by Fraser River sockeye.

In particular, one uncertainty identified by the Commission was whether diseases and pathogens from open net-pen salmon farms in British Columbia contributed to the population decline or may pose risks of significant harm to Fraser River sockeye. In the marine environment of British Columbia, there are 109 licensed salmon farm sites with approximately 60 to 70 farms actively operating at a given time. As net-pen farms reside in the oceanic environment, there exists a potential risk of pathogen exchange between farmed and wild fish. To evaluate the risks and potential impacts of such pathogen transmission events, it is imperative to understand endemic disease effects in both wild and farmed fish species.

Viral hemorrhagic septicemia virus (VHSV) genotype IVa is one endemic pathogen in the marine waters of British Columbia that has spilled over and caused disease in farmed Atlantic salmon *Salmo salar* (Garver et al. 2013). VHSV is common in the Northeastern Pacific Ocean where it has caused disease in numerous marine species, including Pacific herring *Clupea pallasi*, Pacific hake *Merluccius productus*, walleye pollock * Theragra chalcogramma*, and Pacific sardines *Sardinops sagax* (Meyers et al. 1999, Traxler et al. 1999). Laboratory studies conducted on the highly abundant marine species Pacific herring confirmed their extreme susceptibility to viral hemorrhagic septicemia (VHS) disease and demonstrated their capacity to serve as a natural reservoir of VHSV (Kocan et al. 2001, Hershberger et al. 2010). Furthermore, molecular epidemiological investigations suggest that wild Pacific herring and sardines are a source of virus to farmed Atlantic salmon in British Columbia, where VHSV is detected nearly annually (Garver et al. 2013). However, the complete disease ecology of VHSV in the Northeastern Pacific Ocean is uncertain, and the host range of VHSV-IVa undoubtedly extends beyond the aforementioned species. Field surveillance programs have isolated the virus from a number of other marine species, including Pacific salmon species (Amos et al. 1998, Hedrick et al. 2003). The World Organisation for Animal Health (OIE) listed Chinook *O. tshawytscha*, coho *O. kisutch*, and Atlantic salmon as fish species for which there is conclusive evidence of susceptibility to VHSV infection, while sockeye salmon are listed as a species for which there is incomplete data to determine susceptibility (Marty et al. 2006, EFSA 2008, OIE 2019). Laboratory studies exposing freshwater coho, Chinook, pink and sockeye salmon suggest a refractory nature to VHSV (Follett et al. 1997). Yet, given the marine origin of VHSV, it is of interest to evaluate the susceptibility of salmonids to infection at a marine life stage.

To this end, our study investigated the susceptibility of saltwater-phase sockeye salmon to VHSV genotype IVa infection through injection, immersion, and cohabitation exposures.

## 2. MATERIALS AND METHODS

### 2.1. Virus inoculum

Equal concentrations of 4 VHSV genotype IVa isolates were mixed to produce a virus inoculum for each experimental infection. The VHS viruses were isolated from salmonid and non-salmonid species and are representative of the VHSV variants endemic to the marine waters of British Columbia (Table 1) (Garver et al. 2013). The isolates were amplified using the epithelioma papulosum cyprini (EPC) cell line, titered via plaque assay on EPC cell culture (Batts et al. 1991), and diluted in Hanks’ Balanced Salt Solution with 1× Antibiotic-Antimycotic (HBSS+Ab, pH 7.2 to 7.8; GIBCO) to reach the specified exposure levels. At the time of challenge, the virus inoculum was titered to verify that each virus challenge experiment was conducted at the desired exposure level.

### 2.2. Fish

Experimental animals consisted of fish with unknown susceptibility (sockeye salmon *Oncorhynchus nerka*), low to moderate susceptibility (Atlantic salmon *Salmo salar*) and high susceptibility (Pacific herring *Clupea pallasi*) to VHSV. Sockeye salmon fry (Pitt River stock) obtained from a freshwater hatchery

<table>
<thead>
<tr>
<th>Isolate name</th>
<th>Host species</th>
<th>Collection</th>
<th>Location</th>
</tr>
</thead>
<tbody>
<tr>
<td>BC 99-001</td>
<td>Pacific sardine</td>
<td>Jan 1999</td>
<td>Beaver Cove, Vancouver Island</td>
</tr>
<tr>
<td>BC 99-292</td>
<td>Atlantic salmon</td>
<td>Apr 1999</td>
<td>Southern Vancouver Island</td>
</tr>
<tr>
<td>BC 283-1B</td>
<td>Chinook salmon</td>
<td>Feb 2005</td>
<td>West coast of Vancouver Island</td>
</tr>
<tr>
<td>BC 06-089-1</td>
<td>Pacific sardine</td>
<td>Jan 2006</td>
<td>Rennell Sound, Haida Gwaii</td>
</tr>
</tbody>
</table>
were transported to the Pacific Biological Station (PBS, Nanaimo, BC) where they were held in 6°C (±1°C) dechlorinated freshwater under a natural photoperiod. The sockeye remained in fresh water for a minimum of 10 mo and reached average weights of 52 to 80 g prior to saltwater transition and VHSV exposure. Atlantic salmon smolts (Mowi strain) were obtained from a freshwater hatchery and reared at PBS in 10°C (±1°C) brackish water (salinity 6.5) to average weights of 35 to 59 g prior to VHSV exposure. At the time of exposure, the salmon were transitioned to 10°C (±1°C) saltwater (salinity 28 to 30). Specific pathogen-free (SPF) Pacific herring juveniles (average weight: 13 g) reared at the USGS-Marrowstone Marine Field Station (Nordland, WA) were transported to PBS and held in 9.5°C (±0.5°C) saltwater. All fish were fed dry pellets (EWOS) at 1% of their body weight d⁻¹.

2.3. VHSV injection, immersion, and cohabitation exposure of sockeye salmon

The relative susceptibilities of these species to VHSV infection were evaluated by intraperitoneal (IP) injection, immersion, and cohabitation.

For the IP injection (100 µl doses containing 3.9 × 10³ plaque-forming units [PFU] VHSV), VHSV was administered to groups of sockeye salmon (75 fish triplicate tank⁻¹) and Atlantic salmon (72 fish duplicate tank⁻¹). Negative controls for each species consisted of single groups of 75 salmon that were injected with 100 µl of HBSS+Ab in lieu of virus. All treatments were maintained in 450 l tanks supplied with single-pass, UV-treated seawater at 10°C (±1°C). Mortality was monitored daily for up to 21 d post VHSV exposure. Mortalities were collected and immediately stored at −80°C until necropsy and tissue testing for VHSV. Sub-samples of sockeye salmon (10 fish d⁻¹) were sampled from one of the triplicate tanks at 7, 14, and 20 d post VHSV exposure by euthanization in an overdose of buffered TMS; gill, brain, and anterior kidney tissues were collected. The tissue samples were individually flash-frozen in liquid nitrogen and stored at −80°C until RT-qPCR screening for VHSV (Table 2).

Relative susceptibilities of sockeye to VHSV were further evaluated by cohabitation with diseased herring, a highly susceptible host that is capable of shedding substantial quantities of virus (Hershberger et al. 2010). A laboratory-induced VHSV epizootic was initiated, by distributing SPF Pacific herring into each of ten 450 l tanks (63 fish tank⁻¹) supplied with single-pass, UV-treated, 10°C (±1°C) seawater. After a 24 h acclimation period, the water flow was stopped, volume was reduced to 100 l, and VHSV inoculum was added to 8 treatment tanks at a concentration of 7.8 × 10² PFU ml⁻¹. Herring in the remaining 2 tanks served as negative controls and were exposed to HBSS+Ab in lieu of virus. After 1 h, water flow was restored, and tanks were re-filled. Water samples (5 ml) were taken from each tank immediately after inoculation (T₀) and after the 1 h exposure period (T₁h) to verify the waterborne exposure level. The undiluted water samples, along with 1/2 and 1/10 dilutions in Minimal Essential Medium with HEPES, GlutaMAX, Antibiotic-Antimycotic, Gentamycin, and 20% Fetal Bovine Serum (MEM-20+HEPES+Ab, pH 7.6 to 7.8; Gibco), were each plated as 100 µl inoculums on EPC cell culture and monitored for up to 14 d to determine virus titer. The following day, 100 sockeye salmon were added to each of 5 tanks (4 VHSV treatment tanks and 1 negative control tank), and 100 Atlantic salmon were added to each of the remaining 5 tanks. Fish were monitored daily until the experiment was terminated 41 d post exposure; mortalities were collected and stored at −80°C until necropsy and tissue testing for VHSV. The amount of shed VHSV was assessed by collecting 1 l water samples from duplicate tanks in each of the herring + sockeye and herring + Atlantic salmon cohabitant groups at 3, 6, 8, 10, 13, 15, 17, 21, 23 and 28 d post exposure. Viral load in the water samples was determined as described for the T₀ and T₁h sam-
Additionally, 5 fish were sampled from duplicate tanks in each of the herring + sockeye and herring + Atlantic salmon treatments at 6, 10, 15, 21, and 28 d post exposure. In parallel, water and fish were sampled from each of the negative control groups. Sub-sampled fish were euthanized in an overdose of buffered TMS and dissected for gill and anterior kidney tissues. The tissue samples were individually flash-frozen in liquid nitrogen and stored at −80°C until RT-qPCR screening for VHSV (Table 2). To confirm VHSV infection among herring mortalities, the anterior kidney was analysed via virus isolation assay on EPC cell culture.

The potential for sockeye salmon, Atlantic salmon, and Pacific herring to be a reservoir of VHSV was evaluated by comingling survivors from the first cohabitation exposure with naïve sentinels. Surviving fish from the negative control cohabitations were transferred to new tanks in the same arrangement (Table 2). All recipient tanks were disinfected with Ovadine (Syndel) prior to redistribution to prevent contamination. To distinguish the VHSV-surviving herring from sentinel herring, the surviving herring were clipped at the caudal and left pelvic fins. To evaluate VHSV infection status, 5 sentinel and/or 5 VHSV-surviving salmon were sub-sampled from each of the salmon-containing tanks at 19 or 20 d into cohabitation; gill and anterior kidney tissues were collected. The tissue samples were individually flash-frozen in liquid nitrogen and stored at −80°C until RT-qPCR screening for VHSV. Additionally, water samples were collected from the VHSV-surviving Atlantic salmon + sentinel herring tanks on 5, 8, 11, and 12 d into cohabitation. The tanks were monitored daily for up to 42 d, during which mortalities were collected and immediately stored at −80°C (Table 2). To confirm VHSV infection among mortalities, anterior kidney was analysed via virus isolation assay on EPC cell culture.

### 2.4. VHSV detection in tissue samples via RT-qPCR

Total RNA was extracted from fish tissues with TRIzol Reagent (Ambion) following the manufacturer’s protocol. Briefly, tissue samples up to 100 mg were homogenized with a 5 mm stainless steel bead at 25 Hz for 2 min using a TissueLyser (Qiagen). Extracted RNA was suspended in 50 µl RNase- and DNase-free UltraPure Distilled Water (Invitrogen), incubated at 55°C for 10 min, immediately chilled on ice, and stored at −80°C if it was not analyzed on the day of extraction. The concentration and purity of the RNA was determined with an ND-1000 Spectrophotometer (NanoDrop Technologies) with 1.5 µg template being used in a VHSV specific RT-qPCR (Garver

### Table 2. Distribution of sockeye salmon *Oncorhynchus nerka* (ss), Atlantic salmon *Salmo salar* (As), and Pacific herring *Clupea pallasii* (Ph) in VHSV susceptibility studies, including the sampling scheme and duration of each study. For the reservoir potential study, survivors refers to VHSV-exposed fish from the first cohabitation, while sentinels refers to unexposed fish cohabited with the survivors. The + symbol between species indicates a cohabitation; na: not applicable

<table>
<thead>
<tr>
<th>VHSV susceptibility study</th>
<th>Total fish in all replicate tanks</th>
<th>Replicate tanks (n)</th>
<th>RT-qPCR Sampling events (n)</th>
<th>Negative control</th>
<th>Duration (d)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Injection</td>
<td>225</td>
<td>3 ss 2 ss</td>
<td>75 ss 75 ss</td>
<td>20 (ss)</td>
<td></td>
</tr>
<tr>
<td>Immersion</td>
<td>225</td>
<td>3 ss 2 As</td>
<td>75 ss 75 As</td>
<td>20</td>
<td></td>
</tr>
<tr>
<td>First cohabitation</td>
<td>401</td>
<td>4 Ph + ss 3 ss As</td>
<td>64 Ph + 101 ss 64 Ph + 102 As</td>
<td>41</td>
<td></td>
</tr>
<tr>
<td>Reservoir potential cohabitations</td>
<td>196 survivors 130 survivors 129 sentinels</td>
<td>3 ss + Ph 3 As + Ph 2 ss As 1 Ph + Ph</td>
<td>76 ss + 48 Ph 69 As + 28 Ph</td>
<td>39 40</td>
<td></td>
</tr>
<tr>
<td></td>
<td>na 193 survivors 129 sentinels na</td>
<td>1 1 1</td>
<td>na na</td>
<td>na 42</td>
<td></td>
</tr>
<tr>
<td></td>
<td>502</td>
<td>4 Ph + As 5 ss 10 As</td>
<td>64 Ph + 101 As 64 Ph + 102 As</td>
<td>41</td>
<td></td>
</tr>
<tr>
<td></td>
<td>184 sentinels 185 sentinels</td>
<td>1 1 1</td>
<td>69 As + 28 Ph na 69 As + 28 Ph</td>
<td>40 40</td>
<td></td>
</tr>
</tbody>
</table>

*aNegative controls carried on from first cohabitation
*bCohabitation of survivor Pacific herring with sentinel Pacific herring
et al. 2011). Duplicate reactions were run in a Stratagene Mx3000P qPCR System, and data were analyzed using MxPro QPCR Software (Agilent Technologies) with results reported as the ROX normalized average CT value of duplicate reactions. A sample was considered positive for VHSV when a CT value of ≤40 was measured in duplicate reactions. Positive and negative quality controls were included for RNA extraction and at each step of the RT-qPCR assay.

2.5. Statistical analysis

Herring mortality data from the cohabitation study was plotted using Kaplan-Meier survival analysis with Log-Rank testing. Herring survival curves were compared using the Holm-Sidak method of multiple comparisons (SigmaPlot 13.0).

3. RESULTS

Sockeye salmon Oncorhynchus nerka demonstrated nominal susceptibility to VHSV infection after exposure by IP injection. Of the 195 sockeye salmon that were not sub-sampled during the experiment, 1 mortality (0.5%) occurred (Day 17) throughout the 20 d experimental period. Hemorrhaging was observed on the ventral surface of the dead fish, and VHSV was isolated from the anterior kidney at a titer of 9.4 × 10^7 PFU g^−1. RT-qPCR of anterior kidney, gill, and brain tissue confirmed the identity as VHSV. VHSV was detected by RT-qPCR in only 2 of 30 sockeye salmon sub-sampled throughout the study, including 0 of 10 on Day 7, 1 of 10 on Day 14, and 1 of 10 on Day 20 post exposure; average CT values ranged from 19.42 (anterior kidney) to 29.89 (brain). In contrast, mortality was 58% among VHSV-exposed Atlantic salmon Salmo salar (N = 144), and VHSV was isolated from all examined mortalities (n = 10) with mean anterior kidney titers ranging from 1.6 × 10^6 PFU g^−1 to 4.4 × 10^7 PFU g^−1. Mortality did not occur in any of the negative control groups.

Sockeye salmon demonstrated no susceptibility to VHSV infection after waterborne exposure. The actual virus exposure titer determined from tank water samples at the start of challenge was close to the predicted dose, ranging from 2.8 × 10^3 PFU ml^−1 to 5.0 × 10^3 PFU ml^−1, and waterborne VHSV remained detectable in the treatment tanks after 1 h. VHSV was not detected in any water samples from either negative control tank. No mortality occurred among the sockeye salmon or Atlantic salmon that were exposed to waterborne VHSV. Furthermore, VHSV was not detected by RT-qPCR in any of the sockeye salmon sampled on 7, 14, and 20 d post VHSV exposure (n = 10 fish d^−1). No mortalities occurred in the negative control groups.

An active VHSV epizootic in Pacific herring Clupea pallasii was initiated after waterborne exposure to 2.3 × 10^2 PFU ml^−1 to 3.7 × 10^2 PFU ml^−1. After resumption of supply water to the tanks, waterborne VHSV was undetectable at 3 d post exposure but was isolated from the tank water 6 d post exposure and peaked at 10 d post exposure, with an average of 6.0 × 10^2 PFU ml^−1 in the herring + sockeye tanks. Waterborne VHSV reached similar levels across the herring + salmon cohabitant tanks with the exception of 1 tank of herring + Atlantic salmon (Ph + As 1) that had higher levels of virus on Day 10 post exposure (Fig. 1). Survival was significantly lower (p < 0.05) among VHSV-exposed Pacific herring (0 to 27.6%) than among unexposed controls (53.1 to 78.1%) for each cohabitation treatment (Atlantic salmon or sockeye salmon); additionally, survival was significantly lower (p < 0.05) among Pacific herring that were cohabitated with Atlantic salmon than those cohabitated with sockeye salmon (Fig. 2). VHSV-associated mortality began in Pacific herring at 5 d post exposure, and VHSV was recovered on EPC cell culture from 81.5% (n = 54) of the herring mortalities from the sockeye cohabitation and 86.0% (n = 43) of the herring mortalities from the Atlantic salmon cohabitation. VHSV was not de-
mortality occurred among 5.4% (19 of the 355) of Atlantic salmon. Among the Atlantic salmon mortalities, VHSV was detected from 2 of 19 anterior kidney samples via virus isolation assay on EPC cell culture and RT-qPCR, with average C_t values of 34.84 and 36.73. Interestingly, these 2 VHSV-positive Atlantic salmon mortalities occurred in tank Ph + As 1, which yielded 4 of the 19 total Atlantic salmon mortalities and displayed relatively higher levels of shed VHSV (Fig. 1). In contrast, tank Ph + As 2 accounted for 2 of the 19 total Atlantic salmon mortalities with no VHSV detection in anterior kidney. Of the regularly sub-sampled fish, VHSV was transiently detected in gill tissue, but not the anterior kidney, from both sockeye and Atlantic salmon on Days 6, 10, and up to 15 for sockeye salmon, with average C_t values ranging from 32.44 to 37.48. Among negative controls, VHSV was not detected in any sockeye salmon or Atlantic salmon, nor was any virus detected in any Atlantic salmon mortalities (N = 7).

There was no evidence that sockeye or Atlantic salmon served as a reservoir for VHSV. Among 10 tanks containing sockeye and Atlantic salmon that survived prior VHSV exposure, no VHSV was detected from sub-samples of either VHSV-exposed or sentinel salmon. In these tanks, mortality occurred among cohabitating sentinel Pacific herring and Atlantic salmon; however, VHSV was not isolated from the examined mortalities (Table 3) or water samples, and survival of sentinel herring was similar to that of the respective negative controls. In contrast, VHSV was detected in 100% (n = 19) of the sub-sampled mortalities from sentinel herring that were cohabitated with herring that survived prior VHSV exposure (Table 3). VHSV was not detected in any sub-sampled mortalities in the negative control treatments, including herring from the sockeye cohabitation (n = 3), herring from the Atlantic salmon cohabitation (n = 15), or a single Atlantic salmon.

Table 3. Mortality of VHSV-exposed survivors and naïve sentinel fish in the VHSV reservoir potential study. A proportion of the mortalities were tested for VHSV by virus isolation assay (na: not applicable; neg: all samples tested negative; pos: all samples tested positive)

<table>
<thead>
<tr>
<th>Survivors</th>
<th>Cohabitation</th>
<th>Survivors</th>
<th>Sentinels</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Sentinels</td>
<td>% Mortality</td>
<td>Dead fish tested</td>
</tr>
<tr>
<td>Sockeye salmon</td>
<td>Pacific herring</td>
<td>0.0 (0/181)</td>
<td>na</td>
</tr>
<tr>
<td>Atlantic salmon</td>
<td>Pacific herring</td>
<td>3.9 (7/178)</td>
<td>7/7 (neg)</td>
</tr>
<tr>
<td>Sockeye salmon</td>
<td>Atlantic salmon</td>
<td>0.0 (0/120)</td>
<td>na</td>
</tr>
<tr>
<td>Atlantic salmon</td>
<td>Sockeye salmon</td>
<td>4.2 (5/119)</td>
<td>5/5 (neg)</td>
</tr>
<tr>
<td>Pacific herring</td>
<td>Pacific herring</td>
<td>3.3 (2/61)</td>
<td>2/2 (pos)</td>
</tr>
</tbody>
</table>
4. DISCUSSION

Knowledge of the susceptible host species range of a specific pathogen is instrumental for containing its spread and preventing its introduction into pathogen-free areas. Based on criteria established by the World Organisation for Animal Health (OIE), a suspect host species is considered susceptible to infection when a definitive diagnosis of the pathogen occurs from natural transmission events or under experimental conditions mimicking natural pathways of infection (OIE 2013). Herein we provide evidence that saltwater-phase sockeye salmon *Oncorhynchus nerka* fail to fulfill the criteria of a VHSV-susceptible species as no virus was detected or isolated from this species after exposure to the virus under conditions reflective of a natural transmission route. These results extend findings of previous studies that demonstrated sockeye salmon to be refractory to VHSV when exposed in freshwater (Follett et al. 1997, Traxler et al. 1999) and indicate that sockeye salmon likely maintain resistance to VHSV infection through both fresh and seawater phases of their life cycle.

In the Northeast Pacific Ocean, VHSV-IVa is the sole genotype present where it occurs across a broad range of marine fishes (Hedrick et al. 2003, Garver et al. 2013). Pacific herring *Clupea pallasii* is one species that has proven to be a natural reservoir of VHSV whereby under some circumstances, these covert infections cascade to epizootics of VHS disease (Hershberger et al. 2016). During such mortality events, VHSV-diseased Pacific herring shed enormous amounts of virus, resulting in the rapid amplification of exogenous virus that can be a source to sympatric fishes (Hershberger et al. 2013). Such viral transmission events are evident by concurrent isolations of VHSV from Pacific herring and Atlantic salmon *Salmo salar* after aggregations of VHS-diseased Pacific herring were found in and around open net-pen salmon farms of British Columbia (Garver et al. 2013). Given their abundance and contribution as a dominant member of the forage fish assemblage in the North Pacific Ocean, Pacific herring undoubtedly interact with Pacific salmon and have been hypothesized as the source of the uncommon VHSV detections made in Chinook *O. tshawytscha* and coho salmon *O. kisutch* (Meyers & Winton 1995, Amos et al. 1998). Additionally, purse seine and trawl surveys conducted throughout the Salish Sea have documented the co-occurrence and natural interaction of Pacific herring with Pacific salmon species, including sockeye salmon (Grant et al. 2018).

It was not surprising that exposure of Pacific herring, a highly susceptible species, to waterborne VHSV resulted in an epizootic accompanied with copious viral shedding. Levels of waterborne virus shed during the epizootic were relatively equivalent across tanks where the Atlantic and sockeye salmon cohabitants remained free of VHSV. However, in the 1 tank where the sentinel Atlantic salmon became VHSV infected (tank Ph + As 1, Fig. 1), a higher peak level of waterborne virus was observed, suggesting that the infected Atlantic salmon likely shed virus thereby enhancing the level of waterborne virus. Meanwhile, in the tanks where the salmonid cohabitants remained free of VHSV, the levels of shed virus were solely a reflection of the herring epizootic.

The levels of waterborne VHSV generated because of donor herring succumbing to VHS disease reached levels mirroring those measured in natural marine environs. In the vicinity of a high-density Pacific herring spawn-on-kelp fishery, waterborne titers of VHSV were measured at $7 \times 10^2$ PFU ml$^{-1}$ (Hershberger et al. 1999), similar to the peak titer of $6.0 \times 10^2$ PFU ml$^{-1}$ measured in the Pacific herring + sockeye cohabitation tanks. Despite sockeye salmon receiving daily exposure to waterborne virus naturally shed from infected herring, systemic VHSV infections remained absent. However, these cohabitation conditions were sufficient to transmit VHSV to Atlantic salmon, resulting in VHSV infections of the kidney corroborating the susceptibility of this species (Lovy et al. 2013). It is worthwhile noting however that despite the absence of systemic VHSV infections in sockeye salmon exposed to waterborne virus, RT-qPCR detections of VHSV were occasionally made from gill samples. Given that these detections were at similar quantities to tank water samples (K. Garver unpubl. data) and that corresponding kidney tissues remained negative for VHSV, these occurrences were likely a reflection of the presence of virus in the water or superficial association on the surface of the gills rather than an indication of true infection.

Under the exposure conditions employed in these studies, neither sockeye salmon nor Atlantic salmon survivors of prior VHSV exposure were successful reservoirs of the virus. For VHSV-exposed sockeye salmon, the inability to transmit virus to highly susceptible cohabitating Pacific herring is not surprising given that waterborne exposures of sockeye salmon to VHSV failed to initiate an infection. In regard to the reservoir potential of Atlantic salmon, it is unknown why our study failed to demonstrate them as a reservoir species, as transmission of VHSV from previously exposed Atlantic salmon to cohabiting naïve Pacific herring was previously demonstrated (Lovy et al. 2013). Although differences in experimental designs...
preclude direct comparisons between studies, a higher infection rate of Atlantic salmon as reported by Lovy et al. (2013) in comparison to those observed in our study, suggest that the severity of the primary VHSV exposure may be a predicting factor in establishing a reservoir state. It is worth noting that in our study, Pacific herring survivors of prior VHSV exposure successfully transmitted virus to naïve cohabiting herring, revealing their greater propensity as a source of VHSV than either Atlantic salmon or sockeye salmon.

The refractory nature of sockeye salmon to VHSV infection as observed in our controlled laboratory exposure studies is consistent with the absence of a confirmed detection or isolation of VHSV in a natural sockeye salmon population. A presumptive positive RT-PCR detection of VHSV has been reported in juvenile sockeye salmon collected in the marine waters of British Columbia (Marty et al. 2006), whereas surveillance efforts collectively examining >2400 marine-phase sockeye salmon demonstrated freedom of VHSV infection (Kent et al. 1998, Nekouei et al. 2018). The lack of susceptibility of sockeye salmon to VHSV-IVa infections by immersion and cohabitation, together with absence of infection in wild stocks, suggests that waterborne exposure of sockeye salmon will not result in infection.

The mechanism(s) accounting for resistance to VHSV infection are unknown; however, our results indicate that when the virus exposure route circumvented the exterior barrier of the fish via IP injection, virus infection was enhanced. This enhancement of infections, albeit minimal for sockeye salmon (detected in the single mortality and 2 of 30 sub-sampled fish), indicates the importance of the exterior surface of the fish in maintaining the fish’s resistance to VHSV infection. Interestingly, in line with the refractory nature of sockeye salmon as demonstrated in our study, Pacific salmon species in general appear less susceptible to infection with VHSV-IVa, suggesting the potential for shared resistance mechanism(s) among the Pacific salmonids. On the west coast of North America, a limited number of natural detections of VHSV-IVa infections have been reported for Chinook and coho salmon, yet these infections were not associated with VHS disease (Amos et al. 1998). Furthermore, laboratory studies investigating VHSV susceptibility of Pacific salmon have corroborated their limited susceptibility to VHSV-IVa (Winton et al. 1991, Follett et al. 1997, Amos et al. 1998, Traxler et al. 1999, Emmenegger et al. 2013). Nevertheless, the question of whether a fish becomes infected and diseased is undoubtedly contingent upon not only the host species but also the viral strain and environmental conditions under which the virus exposure occurs. This is best exemplified by the fact that marine North American VHSV (genotype IVa) is relatively less virulent to rainbow trout Oncorhynchus mykiss than freshwater European VHSV (genotype I) (Winton et al. 1991, Follett et al. 1997, Snow et al. 2004, Skall et al. 2005, Emmenegger et al. 2013). Consequently, additional studies are required to better understand not only mechanisms of host resistance but also the molecular determinants of VHSV virulence.

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LITERATURE CITED

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