Use of Ultraviolet C (UVC) Radiation to Inactivate Infectious Hematopoietic Necrosis Virus (IHNV) and Viral Hemorrhagic Septicemia Virus (VHSV) in Fish Processing Plant Effluent

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Abstract

We determined the stability of infectious hematopoietic necrosis virus (IHNV) and viral hemorrhagic septicemia virus (VHSV) suspended in either fish processing plant effluent blood water (EBW) or culture media and examined the effectiveness of UVC radiation to inactivate IHNV and VHSV suspended in both solutions. Without exposure to UVC, IHNV and VHSV were maintained in 4°C blood water for up to 48 hours without significant reduction in virus titers. However, when exposed to UVC radiation using a low pressure mercury vapour lamp collimated beam, IHNV and VHSV were inactivated, and the efficacy of UVC radiation was dependent upon the solution and virus type being treated. A 3-log reduction for VHSV and IHNV in culture media was achieved at 3.28 and 3.84 mJ cm−2, respectively. The UV dose needed for a 3-log reduction of VHSV in EBW was 3.82 mJ cm−2. However, exposure of IHNV in EBW to the maximum UVC dose tested (4.0 mJ cm−2) only led to a 2.26-log-reduction. Factors such as particle size, and possible association of viruses with suspended EBW particulate, were not investigated in this study, but may have contributed to the difference in UVC effectiveness. Future work should emphasize improved filtration methods prior to UV treatment of processing plant EBW at an industrial scale.

Keywords: IHNV; VHSV; UVC; Log reduction; Blood water effluent; Fish processing plant

Introduction

Viral and bacterial diseases can affect wild and farmed aquatic animals. In the aquatic environment these infectious agents can spread through the water and infect both commercially harvested and farm-reared fish and shellfish [1]. Disease control in wild and farmed aquatic animals is a complex issue, as the dissemination of infectious diseases in either fresh or salt water can be facilitated by natural current and tidal exchanges as well as through anthropogenic practices such as recreational and commercial boat activities. Additionally, infectious diseases can potentially be introduced from alternate sources such as fish/shellfish processing plants [2]. Consequently, biosecurity measures are among the most effective means to prevent the spread of infectious diseases in the aquatic environment. To mitigate the risk of pathogen dispersion through effluent discharges, alternative disinfection and pathogen neutralization procedures are being investigated.

Current technology used for effluent blood water (EBW) disinfection at fish processing plants is largely dependent on mechanical and/or chemical processes. Processing plants screen effluent for particles larger than 0.5 mm, which are collected and transported off site. However, the remaining EBW, which is high in organics, is generally released into the marine environment for natural dilution to take place [2]. Some fish processors disinfect EBW with chlorine prior to discharging it into the environment; however, organic material is known to inactivate chlorine to varying degrees [3], and many of the plants can not separate organic material from the wastewater or ensure complete mixing of the chlorine and wastewater. Additionally, the higher pH levels of seawater (pH 8) compared to freshwater (pH 6.5 to 7.0) changes the active form of chlorine from hypochlorous acid to hypochlorite which is 100 times less effective as a disinfectant [4] than hypochlorous acid. Therefore, adequate chlorination levels are often not reached and if so are difficult to maintain. Moreover, this type of treatment is labor intensive and leaves room for miscalculation and contamination of the environment with a toxic substance.

An alternative technology for EBW disinfection is Ultraviolet (UV) irradiation. UV irradiation causes inactivation of microorganisms [5] by its effects on DNA and RNA [6]. The UV irradiation is emitted in the UVA, UVB and UVC bands, which are differentiated by the range of wavelength (nanometers). The UVC band emits radiation in a wavelength ranging from 280 to 200 nm, and presents the most germicidal effects [7]. Several studies have examined a variety of microorganisms’ response to UV radiation, and results demonstrated that UV may lead to microorganisms’ inactivation in a time- and dose-dependent manner [7-15]. Although there is no regulatory requirement in North America for treating fish/shellfish processing plant EBW, it is necessary to understand if pathogens remain viable in the EBW, and if so, what disinfectant technologies prove efficacious against potential pathogens. Among the infectious pathogens that are important to monitor for in processing plant EBW are viral agents such as infectious hematopoietic necrosis virus (IHNV) and viral hemorrhagic septicemia virus (VHSV). IHNV and VHSV are among two of the three aquatic rhabdoviruses listed by the OIE (the World Organization for Animal Health). Both virus are widespread throughout the Northern Hemisphere and have caused severe mortality events in cultured and wild fish populations.

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IHNV is the causative agent of infectious hematopoietic necrosis (IHNV) that was first described in sockeye salmon in the 1930s at a hatchery in the Pacific Northwest of USA [16]. Since this initial description the virus has not only been detected in all five species of Pacific salmon but also in rainbow trout/steelhead, Atlantic salmon, and several non-salmonid species [17]. Dependent upon fish species, age, density, water temperature, and virus type, losses during acute HIN outbreaks can reach 95% [18] consequently resulting in devastating economic losses particularly within farmed fish populations. For instance, IHNV epizootics within the Atlantic salmon farming industry in British Columbia, Canada resulted in the estimated loss of $40 million in inventory representing $200 million in lost sales (Odd Grydeland, personal communication).

VHSV, the causative agent of viral hemorrhagic septicemia is considered the most serious disease of farmed rainbow trout in Europe [19]. First described in the early 1930's, VHSV has been detected in an extremely diverse range of fish species in both fresh and seawater environments. Mass mortality events in farmed and wild fish populations have been attributed to this virus and VHS been one of the leading diseases speculated for the collapse and failed recovery of the Pacific herring population in Prince William Sound, Alaska [20-22].

Due to the health threat these viruses pose to cultured and wild fish stocks throughout the Northern Hemisphere, surveillance and control programs are typically undertaken to limit viral distribution. A key component to successful viral disease management programs is in the identification and elimination of potential viral sources. To this end, the objective of this study was twofold: 1) to determine the potential of processing plant effluent to harbour virus by investigating the duration of viability of infectious hematopoietic necrosis virus (IHNV) and viral hemorrhagic septicemia virus (VHSV) in pool effluent at 40C and 15ºC. Tubes were immediately sampled (T-0) by removing a 2 mL aliquot which was subsequently filtered (0.45 µm), serially diluted and plated onto EPC cells. Plates were incubated for 1 hour and then an overlay of MEM-4-HEPEs+Ab was added. Additional sampling was performed at 30 minutes, 3 hours, 6 hours, 24 hours, 48 hours and 7 days post virus seeding of the EBW. Cell culture plates showing CPE (plaque development) were fixed and stained using a crystal violet (0.25 g) and buffered formalin (250 mL) solution.

Similarly, IHNV and VHSV stabilities were determined in culture media (MEM). Media, aliquoted in 15 mL conical tubes, was seeded with 1x10⁶ pfu/mL of either IHNV or VHSV and incubated at 4ºC. Aliquots were removed immediately (T0) as well as at 24, 48, and 72 hours post seeding and virus was quantified using plaque assay as described above.

**UVC radiation studies**

The ultraviolet C (UVC – germicidal) studies were carried out using a collimated beam (Trojan Technologies) equipped with a low pressure mercury vapor lamp (GPH460TSL/4, Trojan Technologies) that produces UVC at 254 nm. UV transmittance (UV intensity transmitted through 1 cm path length of the sample) was determined using a UV spectrophotometer (P245C UV photometer, Trojan technologies). UV intensity at the air-liquid interface was measured using a calibrated radiometer (IL/I700, International Light Technologies).

UVC radiation treatments were applied to two types of solutions: viral culture media (MEM) and fish processing plant EBW. Effluent generated during normal operational loads in a fish processing plant was collected as previously described. Effluent blood water was diluted in distilled water in order to achieve a transmittance of 25%.

Two trials were conducted to determine UV inactivation of IHNV and VHSV when suspended in culture media (trial 1 and 2), while the remaining trial (trial 3) determined the UV inactivation of the two viruses when suspended in EBW. In trials 1 and 2, viruses (IHNV and VHSV) were exposed to 0 and 1.5 mJ/cm² and 0, 1.0, 1.5, 2.0, 2.5, 4.0, 10 mJ/cm², respectively. In trial 3, IHNV and VHSV suspended in EBW, were subjected to UVC doses of: 1.0, 1.5, 2.0, 2.5, and 4.0 mJ/cm². In all studies approximately 1x10⁸ pfu of viruses/mL was added to 25 mL of fish processing plant effluent or culture media. Samples (25 mL) were irradiated in 60 mm x 36 mm sterile Petri dishes. Five minutes prior to and during UV exposure, samples were continuously stirred on a magnetic stir plate at room temperature. The same stir speed was applied to all treatments and all UV exposure trials were carried out using the same batch of effluent blood water or media.

**Evaluation of UVC radiation treatments**

The efficacy of the UVC sterilization treatments were evaluated by comparing quantities of infectious virus pre and post UVC treatments.
Infectious virus was quantified using the plaque assay [23]. The dose response curves of the effects of UVC irradiation on IHNV and VHSV infectivity are presented as log10-reduction values, which were calculated relative to dose zero. The log-reduction data was subjected to the least-square linear model using Sigma-Stat (Systat Software Inc, Chicago IL), and data was checked for normality (Shapiro-Wilk) and constant variance (Systat Software Inc, Chicago IL). Log-reduction values obtained from replicate well counts are presented in the graphs. The stability curve data is also presented as log10 values (mean ± SEM) obtained from replicate well counts and ANOVA (Systat Software Inc) was performed to determine differences in viruses’ titration during 3 days.

**Results**

**Virus stability in processing plant effluent versus culture media**

IHNV and VHSV were less stable (more rapidly inactivated) when suspended in 15°C effluent blood water than when suspended in either culture media or EBW at 4°C (compare Figures 1A, B and C). At 48 hours post inoculation, no IHNV or VHSV was detected in 15°C blood water despite being seeded at 4x10^5 PFU/mL and 1x10^6 PFU/mL, respectively (Figure 1B). Conversely, viral titres for IHNV and VHSV suspended in EBW or culture media or at 4°C remained fairly consistent out to 48 hour post inoculation (Figures 1A & C). In comparing the stabilities of VHSV and IHNV in EBW, it was revealed that VHSV was inactivated faster than IHNV at both 4°C and 15°C (Figures 1A & B). Viral titres for IHNV suspended in EBW at 4°C remained unchanged up to 48 hours post inoculation while VHSV titers were reduced by one log (Figure 1A).

**UVC inactivation of IHNV and VHSV**

Trial 1, in which IHNV and VHSV suspended in culture media were subjected to UVC irradiation at a dose of 1.5 mJ cm^-2 resulted in a 1.72 and 2.05 log reduction in IHNV and VHSV, respectively (data not shown). In trial 2, exposure of IHNV and VHSV in culture media to UVC at a dose of 1.5 mJ cm^-2 led to a 1.43 and 1.83 log reduction, respectively (Figures 2A and 3A).

The results in Figure 2A showed that the UV dose needed for a 3-log reduction of IHNV suspended in culture media was 3.84 mJ cm^-2.
Exposure of IHNV to 10 mJ cm\(^{-2}\) resulted in a 7.1-log reduction. Based on the least squares linear regression model equation, a UV dose of 8.29 mJ cm\(^{-2}\) is required for 6-log reduction. For IHNV suspended in EBW, a 2.26-log reduction was obtained at UV dose of 4 mJ cm\(^{-2}\) (Figure 2B).

The UV dose needed for a 3-log reduction of VHSV suspended in culture media (Figure 3A) was 3.28 mJ cm\(^{-2}\). Exposure of VHSV to 10 mJ cm\(^{-2}\) resulted in a 7.4-log reduction. Based on the least squares linear regression model equation, UV dose of 7.87 mJ cm\(^{-2}\) is required for a 6-log reduction. For VHSV suspended in EBW, a log reduction of 3.12 was obtained at UV dose of 4 mJ cm\(^{-2}\) (Figure 3B). The UV dose required for a 3-log reduction is 3.82 mJ cm\(^{-2}\).

**Discussion**

In this study we determined the stability of IHNV and VHSV in effluent from a fish processing plant as well as the effectiveness of UVC irradiation in sterilizing such wastewater. The aquatic rhabdoviruses, IHNV and VHSV, although exhibited reduced viability in fish processing effluent in comparison to the control media, did persist in blood water (either at 4 or 15°C ) for a period of six hours without significant reduction in titers. Viable virus was also found at longer time points post inoculation, however, titers did begin to decline with the rate being dependent upon temperature and species of virus. Due to the ability of IHNV and VHSV to persist in fish/shellfish processing plant effluent for several hours and the fact that wastewater is discharged quickly from plants, there is a risk that infectious viruses could be transferred into open environments via untreated effluent releases when infected fish are being processed.

Using laboratory scale low pressure UV collimated beam, we examined the effectiveness of UVC irradiation for inactivating IHNV and VHSV in fish processing plant effluent blood water (EBW). It is noteworthy that for both IHNV and VHSV, a 1.46 and 1.16 fold lower UV dose, respectively was required in culture media versus blood water to achieve equivalent virus inactivation. Thus, the effectiveness of UVC irradiation was greater in culture median than in effluent blood water from fish processing plants. These results concur with other studies that investigated viral and bacterial reduction in UV treatment systems applied to different types of effluents [7,11]. It is possible that the higher UVC doses required in blood water to achieve equivalent inactivation values for VHSV and IHNV suspended in culture media is due to the presence of particulate matter which affects the transmittance of UVC rays in the solution [24]. Among other particulate matter, processing plant EBW can contain varying levels of soluble proteins, fats, oils, and bacteria [2]. It has been documented that particulate matter in effluents can cause a shielding effect, which protects organisms against the UV radiation [25]. Additionally, UV radiation effectiveness has been shown to be dependent upon the type of particular matter (chemical or biological) [26]. Particulate material can also facilitate virus transfer. It has been reported [27] that IHNV can adhere to different particulate material which subsequently can serve as a source of virus infection. These authors also showed that the adsorption rate of IHNV to different suspended solids was not affected by the solution's pH (acid, neutral, and alkaline). Thus, to maximize UVC effectiveness in disinfecting fish/shellfish processing plant effluent, steps should be taken to further remove particulate materials prior to UV treatment.

In addition to utilizing different UVC doses for the solution being treated (i.e. wastewater vs. culture media) it was observed that UVC dose requirements may vary dependent upon the virus type being treated against. Regardless of whether VHSV was suspended in culture media or effluent blood water, a lower UV dose was required to obtain equivalent inactivation levels as those observed for IHNV. For instance, a UVC dose of 3.28 mJ cm\(^{-2}\) was required for a 3-log10 reduction of VHSV when suspended in culture media as compared to 3.84 mJ cm\(^{-2}\) for an equivalent reduction of IHNV in culture media. Differing UV sensitivities have been observed between infectious pancreatic necrosis virus (IPNV), infectious salmon anemia virus (ISAV), and VHSV with both IPNV and ISAV exhibiting higher tolerance to UVC than VHS virus. When suspended in freshwater, VHSV, ISAV, and IPNV presented a 3-log reduction a UV dose of 3.3, 7.9 and 11.88 mJ cm\(^{-2}\), respectively [7]. It has been suggested that size and type of virus acid nucleic genome could affect UV resistance, where single-stranded nucleic acids viruses (such as VHSV and IHNV) are less resistant to UV radiation than double-stranded nucleic acids [7]. Due to similarity of IHNV and VHSV, the genome type and size may not be the only factors influencing viruses’ resistance to UV radiation. A recent study that has examined the effects of UVC radiation in RNA (Polivirus 1, Cosakievirus B4, and bacteriophage MS2 - all single stranded linear RNA) and DNA (Adenovirus, Myoviridae, Siphoviridae, etc - all double stranded linear DNA) viruses have shown that the size and type of virus or its nucleic acid genomes could not be used to predict UV radiation effectiveness [13]. The authors suggested that the sensitivity of viruses to UVC treatments may be related to chemical and physical differences in the virion.

Although it is difficult to compare UVC doses across studies due to differences in methodologies, our UVC inactivation rates reported herein corroborate with those reported by Øye and Rimstad [7]. In our study, the UVC dose needed for a 3-log reduction of VHSV suspended in culture media was 3.28 while Øye and Rimstad [7] observed that...
3.30 m2 was necessary for a 3-log reduction of VHSV suspended in freshwater. On the other hand, VHSV suspended in phosphate buffer saline (PBS) with coffee (0.4% v/v) required only 1.8 4 m2 to achieve a 3-log reduction [14]. Differences in UVC inactivation rates between studies demonstrate the importance of having standardized methodologies to test the effectiveness of UVC treatments for viruses in different suspension solutions.

In summary, we demonstrated that both IHNV and VHSV are stable in EBW, but their survivability is affected by both time and temperature. In addition, the results revealed that VHSV seems to be less resilient than IHNV when suspended in EBW. Both viruses were also sensitive to UVC treatment, but the efficiency of the UVC treatment was more pronounced in culture media than in the EBW. Factors such as particle size, and possible association of viruses with EBW particulate, were not investigated in this study, may have contributed to the difference in UVC effectiveness. Although the UVC system showed potential to inactivate viral microorganisms at laboratory scale experiments, it will be important to demonstrate whether a commercial scale UVC system would be efficient for total microorganisms’ inactivation. Thus, not only dose (energy required to inactivate microorganisms) but also the inclusion of improved filtration systems prior to UVC treatments requires further investigation.

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