The impact of aquaculture soundscapes on whiteleg shrimp *Litopenaeus vannamei* and Atlantic salmon *Salmo salar*

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ABSTRACT: Research into the effects of soundscapes on aquaculture species in key production systems is sparse, despite potential impacts of sound on animal welfare and commercial yields. In the following study, 2 high-value global aquaculture species, whiteleg shrimp *Litopenaeus vannamei* and Atlantic salmon *Salmo salar*, were exposed to aquaculture production system soundscapes. For shrimp, sound recordings of a commercial recirculating aquaculture system (RAS) were played back at a sound pressure level (SPL) of 128 dB re 1 µPa, and for salmon, recordings from a commercial sea pen production system were played back at an SPL of 127 dB re 1 µPa for an 8 wk period. Effects of exposure on growth, survival, and indications of metabolic stress were measured as parameters of interest for aquaculture production. Mean growth performance and survival rates did not differ significantly between sound and control treatments for either species. Blood and haemolymph parameters from both species indicated no measurable change in metabolic status or stress levels. Slight, but non-significant, increases in total haemocyte count and, in particular, hyaline cell count were recorded in shrimp exposed to sound. Slight, but non-significant reductions in overall weight gain were recorded in sound-exposed salmon. Overall, the results indicate that sound exposure in current production systems does not negatively affect the early grow-out stage of these key species, either due to rapid habituation or higher hearing thresholds of hatchery-produced individuals, and that no measurable stress response occurs in sound-exposed animals. In future studies, response of the studied species to acute sound exposure and the response of earlier, and potentially more sensitive, life-stages will need to be determined to ensure optimal welfare and production performance.

KEY WORDS: Noise · Stressor · RAS · Sea pen · Pond

1. INTRODUCTION

Understanding and managing the effects of multiple stressors in aquaculture systems is central to ensuring efficient and humane production (Ashley, 2007, Lucas & Southgate 2012). Anthropogenic sources of underwater sound are potentially high-impact stressors in aquaculture environments and cannot be avoided by cultured animals (Popper 2003). Soundscapes in aquaculture systems have been shown to exceed both the physiological and behaviourally determined hearing thresholds of numerous vertebrate and invertebrate aquaculture species (Bart et al. 2001, Ladich & Fay 2013, Radford
& Slater 2019). Despite this, the growth, immune and stress response of key species to sound exposure in various commercial aquaculture production systems remains poorly investigated.

The acoustic habitats prevailing in recirculating aquaculture systems (RAS) and sea pens are highly variable, and the majority of the sound energy falls within the 100 to 500 Hz range, at or above fish hearing thresholds (Bart et al. 2001, Craven et al. 2009, Radford & Slater, 2019). Radford & Slater (2019) showed that RAS and sea pen sound levels exceed the hearing thresholds of the commonly cultured fish common carp *Cyprinus carpio* (Popper 1972), Atlantic salmon *Salmo salar* (Hawkins & Johnstone 1978) and European perch *Perca perca* (Wolff 1967), while the sound levels of aquaculture pond systems exceed the hearing thresholds of common carp (Amoser & Ladich 2005). Effects of long-term exposure, relevant to outgrowing aquaculture are unknown.

Sound response in non-mammalian aquatic species has primarily been investigated in short-term exposure experiments playing back extreme anthropogenic noise levels exceeding 170 dB re 1 µPa (e.g. from pile-driving, boats and air guns [seismic surveys]). These have been shown to alter fish and crustacean (whiteleg shrimp *Litopenaeus vannamei* and common shrimp *Palaemon serratus*) behavior and physiology, i.e. avoidance (Feist 1991, Sand et al. 2000, Filiciotto et al. 2016, Zhou et al. 2016), and to decrease fish growth (Filiciotto et al. 2013), severely damage hearing capacity (McCauley et al. 2003) and increase physiological indicators of stress (Wysocki et al. 2006). The effects may impact aquaculture production efficiency. More recently, playbacks of offshore soundscapes (boat noise) and onshore (concrete tank) aquaculture environments have been shown to significantly affect oxidative status and immune stress indicators in juvenile sea bream *Sparus auratus* (Francesco et al. 2017). However, animals may acclimatise to sound exposure within weeks or months, as has been shown with repeated playback of impulsive sound to European seabass *Dicentrarchus labrax* (Neo et al. 2018) and ship noise to threespot dascyllus *Dascillus trimaculatus* (Ned-elec et al. 2016).

In the present study, sea pens and RAS soundscapes were chosen for exposure playback experiments. Sea pen systems dominate marine finfish production worldwide totaling >5 million metric tonnes for Atlantic salmon, European seabass, sea bream and Atlantic bluefin tuna *Thunnus thynnus*. Commercial use of RAS systems is rapidly expanding for the production of early stages of many finfish species, along with full-cycle production of salmonids, other high-value finfish and shrimp species (Joenson 2016, Slater 2018).

Animals are exposed to single or a series of soundscapes throughout their production in aquaculture, and acoustic habitats may influence animal behaviour, metabolic status or immune status. This could lead to changes in feeding success, growth performance, immune response and even survival. In the present study, passive acoustic recordings from offshore sea cages and onshore RAS were played back in controlled holding tanks at relatively high sound levels to Atlantic salmon and whiteleg shrimp, 2 aquaculture species of very high commercial value. Parameters applied to measure aquaculture performance and animal welfare, growth, survival and standard measures of metabolic and immune response were measured after an 8 wk experimental period.

2. MATERIALS AND METHODS

All methods applied were approved by the ethics officer of the Alfred Wegener Institute and registered with the Senator of Health’s Veterinary Authority Bremen, Germany, in accordance with the EEC Directive 86/609 and EU Directive 2010/63/EU.

2.1. Shrimp system

Postlarvae whiteleg shrimp *Litopenaeus vannamei* were sourced from a commercial hatchery in Florida, USA and on-grown in a RAS system at the Alfred Wegener Institute’s (AWI) Center for Aquaculture Research in Bremerhaven, Germany. Water salinity in the holding system was maintained at 20 psu and temperature at 28.5°C until inclusion in the experiment. Juvenile shrimp were weighed to the nearest 0.01 g after excess water was removed by blotting. The shrimp were allocated to 6 groups of 50 individuals with a mean (±1 SD) weight of 5.79 ± 1.54 g (min. group mean: 5.58 ± 1.67 g, max. group mean 6.18 ± 1.49 g).

The groups of 50 shrimps were randomly allocated to 6 experimental tanks (dimensions 1.0 m × 1.0 m × 0.7 m, L × W × H, water depth of 0.6 m) and supplied with water at a rate of 10 l min⁻¹ (complete water exchange time, ca. 1 h). The natural light cycle was maintained throughout the experiment. The specially designed RAS consisted of the 6 freestanding 600 l tanks combined with a RAS water treatment
system maintained 3 m from the closest tanks (total system volume 3.8 m³) including a mechanical solids filter, nitrification biofilter and UV sterilisation. Physical water parameters (oxygen, pH, temperature and salinity) were measured daily. Concentrations of \( \text{NH}_3/\text{NH}_4^+ \), \( \text{NO}_3^- \) and \( \text{NO}_2^- \), were determined twice per week using a QuAAtro 39 Continuous Seg-mented Flow Analyzer (SEAL Analytical). Animals were fed Crevetec Grower 1A sinking commercial shrimp diet (CreveTec), at a daily feeding ration of ~3% of shrimp bodyweight as calculated and adjusted weekly following standard growth tables for whiteleg shrimp.

The 6 experimental tanks were divided into 2 sub-systems consisting of 2 parallel rows of 3 tanks. All inflow pipes and outflow pipes were laid separately from the pump manifold to individual tanks and back to the sump to avoid sound transfer between tanks. Water inflows were submerged below the tank water surface to reduce inflow sound disturbance. All pumping, water purification and aeration systems were separated from the tank system by 3 m PVC piping (Ø 150 mm) and all potential tank sound sources were excluded. In all 6 tanks, a single UW30 underwater speaker (Lubell Labs, USA) was hung in the tank center at 25 cm below the water surface. In the 3 ‘exposure’ tanks, speakers were connected to an 800 W 12V 4 channel amplification system (Mac Audio, Germany) and (continuous) playback of WAV files was conducted with a commercial MP3 / WAV audio player (ClipSport, SanDisk, USA) through the amplification system with the same series of looped snippets played to each separate exposure tank as per methods described by Filiciotto et al. (2013, 2016).

### 2.2. Shrimp sound exposure

Radford & Slater (2019) recorded 72 h of RAS noise from 3 locations (close to pump house; middle of tank; close to office) from 0.7 m deep concrete tanks at the Garnelen Farm Grevesmühlen, Germany. One hundred random 20 s broadband (0.02 to 24 kHz) ‘snippets’ were selected from the 3 hydrophone recordings and looped for playback using multiple recordings over different time frames and days as suggested by Slabbekoorn & Bouton (2008). Play-backs were calibrated using ST300 hydrophone (Oceaninstruments; flat frequency response 20 Hz to 60 kHz) at 48 kHz giving an upper analyzable frequency limit of 24 kHz. All hydrophones were calibrated using a sound calibrator (G.R.A.S. Type 42AB) producing a 1 kHz signal at 114 dB re 1 μPa. Full bandwidth (50 Hz to 24 kHz) sound pressure level (SPL) was adjusted to 128 dB re 1 μPa in treatment tanks and maintained for 8 wk. This level was selected to represent maximum root mean squared (RMS) levels measured at the shrimp farm over a 24 h period (Radford & Slater 2019). In control tanks, speakers were not connected to a sound-producing device and sound level in control tanks was, however, determined as for exposure tanks. Recorded sound level in control tanks was 95 dB re 1 μPa (Fig. 1A).

### 2.3. Shrimp sampling

After 8 wk, all animals were harvested and placed into an ice slurry for further analysis, weighed to the nearest 0.01 g and length measured to the nearest 0.1 mm. Haemolymph was extracted at harvest individually from 3 animals per tank (9 samples per treatment) to determine total haemocyte count (THC), haemocyte type and phenoloxidase activity. Syringes were filled with 100 μl of sterile EDTA-free buffer (27 mM trisodium citrate, 385 mM sodium chloride, 115 mM glucose, pH 7.5) (Huang et al. 2010), and an equal amount of haemolymph was withdrawn (1:2 dilution) from the ventral sinus. Individual samples were kept on ice before they were centrifuged at 4°C and 800 × \( g \) for 10 min. Supernatant was frozen at −80°C until further analysis. To avoid pyrogen effects in phenoloxidase activity determination, all consumables were autoclaved.

Phenoloxidase was determined by formation of dopachrome from L-DOPA analyzed kinetically using an Infinite 200 PRO spectrophotometer (Tecan) applying technical triplicate measurement. Activity measurements were performed after Huang et al. (2010), modified for a 96 well microtiter plate with 10 μl diluted plasma sample added to 215 μl L-DOPA solution (3 mg ml⁻¹ in 0.1 M potassium phosphate buffer, pH 6.6). Absorption at 490 nm was recorded every 20 s for 200 s. All measured activities were plotted against time, and replicates were excluded from analysis where activity development over the measurement period tended to the non-linear (linear regression \( r^2 < 0.8 \)). One unit of enzyme activity (U) is defined as a linear increase in absorbance of 0.001 per min per ml haemolymph.

Total haemocyte count (THC) and differential haemocyte count (DHC) were determined by flow cytometric analysis after Owens & O’Neill (1997)
using a BD Accuri C6 (Becton-Dickinson). Samples were analyzed using forward scatter (FSC), proportional to cell size, and side-scatter (SSC), correlating with granularity or interior structure of the cell. Between 30,000 and 120,000 events were recorded per sample, depending on the cell density in the sample and the cell sub-population dispersal. Before measurement, haemocyte samples were filtered through a 45 µm syringe filter (to exclude debris). Scatter plots (FSC vs. SSC) were generated for each sample with FSC data at linear and SSC data at log scale (Xian et al. 2009). Grading was performed using the BD Accuri C6 software (v.1.0 2011) to define haemocyte cell types and exclude unwanted events by visual inspection of distinct event clusters in the dot plots. Haemocytes could be divided into 3 sub-populations based on size (hyaline, semigranular and granular cells). The total haemocyte count was obtained by summing the number of cells per ml of the 3 population subsets (from the DHC) and values were expressed as the number of haemocytes \( \times 10^5 \) ml\(^{-1} \).

### 2.4. Salmon system

Atlantic salmon *Salmo salar* parr were sourced from a commercial hatchery in Hirtshals, Denmark, at an average weight of ~100 g and immediately introduced to full saltwater conditions (30 psu and 14.5°C) in a RAS on arrival at AWI. The animals were held in these conditions for a period of 7 wk in tanks (dimensions 2.0 x 2.0 x 0.8 m L x W x H, water depth of 0.6 m) prior to experimental onset. The holding tank conditions (pump and filter position, inflows, physicochemical water parameters) and prevailing soundscape in the holding system did not differ from the level in the control tanks. At experimental onset, salmon were anaesthetised with MS-222 (~100 mg l\(^{-1}\) tricaine mesylate), weighed to the nearest 0.1 g and measured to the nearest 0.5 cm length. The salmon were allocated to 6 groups with 20 individuals. Mean (±1 SD) weight was 177.9 ± 34.2 g across all groups (min. 175.0 ± 29.4 g, max. 181.1 ± 39.3 g). At experimental onset, 5 individuals of the total remaining salmon that were not assigned to a group were

![Figure 1](image_url). Mean power spectral densities (dB re 1 µPa² Hz\(^{-1}\)) for ambient sound (blue line), playback sound (red line) and source sound (green line) for (A) whiteleg shrimp *Litopenaeus vannamei* and (B) Atlantic salmon *Salmo salar* sound exposure.
anaesthetised and euthanised with an excess dosage of MS-222 500 mg l\(^{-1}\), and caudal vein blood was collected for further analysis.

The groups of 20 individuals were randomly allocated to the 6 experimental tanks as described above (dimensions 1.0 × 1.0 × 0.7 m L × W × H, water depth of 0.6 m) supplied with water at a rate of 20 l min\(^{-1}\) (complete water exchange time ca. 30 min). The light cycle was maintained at 12 h light:12 h dark throughout the experiment; water was maintained at 14.5°C, and salinity at 30 psu.

2.5. Salmon exposure

Radford & Slater (2019) recorded 144 h of sea pen noise from 3 locations (middle of sea pen and 2 locations on the outside of the sea pen) from NZ King Salmon (farm located at Ruakaka Bay, Marlborough Sounds, New Zealand). The farm consists of 12 sea cages with a surface area of 225 to 250 m\(^2\) and a depth of 15 m mounted on a floating barge and floating high-density polyethylene (HDPE) walkway system. Cages were stocked at varying densities and sizes of chinook salmon *Oncorhynchus tshawytscha*. In addition to the floating structures and the stocked fish, potential sound sources include access vessels, an associated feed barge, oxygen storage, compressed air automated feeders and a temporary handling facility linked to the site. One hundred random 20 s broadband (0.02 to 24 kHz) ‘snippets’ were selected from the 3 hydrophone recordings and looped for playback. Playbacks were calibrated using ST300 hydrophone (Oceaninstruments; flat frequency response 20 Hz to 60 kHz) giving an upper analyzable frequency limit of 24 kHz. All hydrophones were calibrated using a sound calibrator (G.R.A.S. Type 42AB) producing a 1 kHz signal at 114 dB re 1 µPa. SPL was adjusted to 127 dB re 1 µPa in treatment tanks and maintained for 8 wk. This level was selected to represent maximum RMS levels measured at the salmon farm. In control tanks, speakers were not connected to a sound-producing device and sound level in control tanks was, however, determined as for exposure tanks. Recorded ambient sound level in control tanks was 96 dB re 1 µPa (Fig. 1B).

2.6. Salmon sampling

Animals were fed twice per day with sinking commercial salmonid feed (Skretting Optiline F1-P and F2-P) ad libitum, ~3% bodyweight as calculated by standard growth tables for Atlantic salmon. After 8 wk, 3 animals per replicate were anaesthetised and euthanised with an excess dosage of MS-222 500 mg l\(^{-1}\), weighed (±1 g), and measured (length, ±0.5 cm). Blood was collected from the caudal vein and liver weight (±1 g) determined to calculate the hepatosomatic index (HSI). All remaining animals were anaesthetised with 100 mg l\(^{-1}\), weighed to the nearest gram and measured to the nearest 0.5 cm length.

Body indices weight and length were used to calculate weight gain (G, in g), specific growth rate (SGR, as % daily body weight gain) and HSI as follows:

\[
G = \text{weight}_{\text{End}} - \text{weight}_{\text{Start}} \tag{1}
\]

\[
SGR = 100 \times \frac{\ln \text{weight}_{\text{End}} - \ln \text{weight}_{\text{Start}}}{d} \tag{2}
\]

\[
\text{HSI} = \frac{\text{Liver weight}}{\text{total body weight}} \tag{3}
\]

\[
K \text{ (Condition Factor)} = 100 \times \left(\frac{\text{total body weight}}{\text{total body length}^3}\right) \tag{4}
\]

2.7. Blood analyses

A subsample of 0.5 ml blood from each fish was used for determination of pH using a IntellicalTM PHC 108 pH sensor (Hach) and blood lactate content to the nearest 0.1 mmol l\(^{-1}\) using a Gemar Lactate Scout automated lactate analyser (Gemar). A second subsample was transferred to capillary tubes H7 812 (Servoprax) to determine haematocrit using a SERVOspin HKT TC-12K haematocrit centrifuge (Servoprax). Samples were spun at 7700 × g for 15 min and separated haematocrit determined by measuring fraction length and calculating percentage of total. Remaining blood was centrifuged at 2000 × g at 4°C to separate the plasma. Plasma glucose was determined by a Fuji Dry Chem NX500i blood autoanalyser with Fuji Dri-Chem Slides GLU-PIII (Fujifilm Europe). Plasma cortisol was determined after Reiser et al. (2010) as adapted by Fuchs et al. (2017) using enzyme-linked immunosorbent assays (ELISA) (RE52611, IBL International). A 100 µl blood plasma aliquot was denatured at 80°C for 1 h, vortex agitated for 20 s, diluted in 0.05 M PBS and vortexed for 20 s more. Samples were centrifuged at 13 000 × g for 20 min before a 50 µl sample of supernatant was used for the ELISA in duplicates, and photometric measurement was conducted with a microtitre plate reader (TriStar LB941 Berthold Technologies).
2.8. Statistics

Growth performance and blood/haemolymph parameter data were analysed for normality (Shapiro-Wilk's test) and homoscedasticity (Levene test). The individual sample values within tanks which followed a normal distribution and exhibited homoscedasticity were further compared to control values using a 1-way analysis of variance (ANOVA) with a nested design with tanks nested within treatments. In the case of shrimp and salmon growth and length measurements where varying tank n-values excluded the use of a nested design, a 2-way ANOVA design was applied with a forced Tukey's post-hoc analysis of tanks nested within treatment (noise and control) to test for tank effect in a pairwise manner. SGR was calculated as a whole tank value (as individuals were not tagged) and data tested using a 1-way ANOVA. All statistical analyses were performed using SigmaPlot Software v. 11.0 and Real-Statistics Software templates for Excel 2019.

3. RESULTS

3.1. Sound playback

The ambient noise (in control tanks) in both the shrimp *Litopenaeus vannamei* and salmon *Salmo salar* playback RAS was low, and the shrimp and sound playback significantly increased the sound within the treatment tanks across the entire frequency range (Fig. 1). The playback sound for shrimp was missing a low frequency peak at 200 Hz; the rest of the bandwidth closely matched the source file (Fig. 1A). Playback for the salmon treatment closely aligned with the source file for each treatment (Fig. 1B).

3.2. Shrimp growth and survival

Shrimp survival across treatments and controls was 67 ± 9% (mean ± SD) with survival slightly lower in control tanks (65 ± 9%), as opposed to sound treatment (69 ± 9%). At experimental completion, shrimps in the sound treatment weighed 17.3 ± 0.9 g and were 14.0 ± 0.2 cm in length. Shrimp from the control treatment exhibited very similar growth rates, weighing 16.9 ± 1.3 g and measuring 13.8 ± 0.3 cm (Fig. 2). Shrimp weight and length did not differ significantly between control and sound exposure treatments ($F_{1,195} = 1.095, p = 0.297$ and $F_{1,195} = 2.056, p = 0.153$, respectively). Pairwise comparison of tanks within treatment and controls revealed no significant tank effect for shrimp weight with the greatest mean difference within controls at Tank 3 vs. Tank 2 ($N = 3; Q = 3.246, p = 0.056$) and within sound treatments at Tank 1 vs. 2 ($N = 3; Q = 3.217, p = 0.059$), within sound treatments is ($N = 3; Q = 2.612, p = 0.154$).

3.3. Shrimp immune and stress response

Activity of phenoloxidase was highly variable across all measurements and treatments. A total of 21 technical replicates in the control and 24 technical replicates in the sound treatment were analyzable for activity; means were calculated from technical replicates to provide values corresponding to individual phenoloxidase activity of 7 control and 9 sound-exposed animals, respectively. Activity was higher amongst shrimps in the control (287.9 ± 113.8 U ml$^{-1}$, mean ± SD) than for those in the sound treatment (184.3 ± 116.3 U ml$^{-1}$), but values did not differ significantly across treatments ($F_{1,15} = 1.669, p = 0.265$) with no tank effect (Tank [Treatment] $F_{4,16} = 1.632, p = 0.246$; Fig. 3).

3.4. Shrimp total haemocyte count (THC) and differential haemocyte count (DHC)

Total haemocyte counts differed between control and sound-exposure treatment with animals from the sound-exposure treatment (241.57 ± 34.1) exhibiting...
higher values than the controls (165.40 ± 73.1), although differences between groups were not statistically significant \( (F_{1,17} = 1.154, p = 0.343) \) with no tank effect \( (\text{Tank \ [Treatment]} \ F_{4,17} = 2.248, p = 0.124) \).

The differential fractions of counted haemocytes differed between treatments, with the sound treatment \( (29.88 \times 10^5 \pm 2.3 \times 10^5 \text{ cells ml}^{-1}) \) exhibiting a higher overall first fraction (estimated hyaline cell fraction) in absolute cell count compared to the control \( (25.50 \times 10^5 \pm 3.2 \times 10^5 \text{ cells ml}^{-1}) \). However, differences were not significant \( (F_{1,17} = 0.077, p = 0.794) \) with no tank effect \( (\text{Tank \ [Treatment]} \ F_{4,17} = 0.367, p = 0.827) \).

In the second fraction of larger cells (estimated semi-granular and granular cell count), the sound-exposure treatment exhibited higher cell counts \( (211.7 \times 10^5 \pm 32.5 \times 10^5 \text{ cells ml}^{-1}) \) than control animals \( (139.9 \times 10^5 \pm 64.9 \times 10^5 \text{ cells ml}^{-1}) \). However, results did not differ significantly \( (F_{1,17} = 1.296, p = 0.318) \) between treatment and control, and no tank effect was detected \( (\text{Tank \ [Treatment]} \ F_{4,17} = 0.121, p = 0.0.121) \) (Fig. 4).

### 3.5. Salmon growth and survival

Salmon survival was similar between controls \( (98 \pm 3\%) \) and sound treatments \( (95 \pm 5\% \text{ mean} \pm \text{SD}) \). Salmon final length was also similar between treatments (sound: \( 29.2 \text{ cm} \pm 0.1 \text{ mean} \pm \text{SD} \); control: \( 29.3 \pm 0.2 \text{ cm} \)) \( (F_{1,115} = 0.212, p = 0.646) \). Pairwise comparison of tanks within treatment and controls revealed no significant tank effect for salmon length with the greatest mean difference within controls in Tank 1 vs. Tank 3 \( (N = 3; Q = 1.494, p = 0.543) \) and within sound treatments in Tank 1 vs. 2 \( (N = 3; Q = 1.987, p = 0.342) \).

Mean \( (\pm 1 \text{ SD}) \) final salmon weight in sound treatments \( (277.9 \pm 9.7 \text{ g}) \) was slightly lower than in the controls \( (290.2 \pm 12.5 \text{ g}) \) but was not significantly different \( (F_{1,110} = 1.632, p = 0.204) \). Pairwise comparison of tanks within treatment and controls revealed no significant tank effect for salmon weight, with the greatest mean difference within controls in Tank 2 vs. Tank 1 \( (N = 3; Q = 2.561, p = 0.171) \) and within sound treatments in Tank 1 vs. 3 \( (N = 3; Q = 1.987, p = 0.342) \) (Fig. 5).

Mean final salmon condition factor \( (K) \) was higher in control tanks \( (1.14 \pm 0.03) \) than in sound treatments \( (1.11 \pm 0.03) \) but values were not significantly different \( (F_{1,110} = 3.329, p = 0.071) \). Pairwise comparison of tanks within treatment and controls revealed no significant tank effect for \( K \) with the greatest mean difference within controls in Tank 1 vs. Tank 3.
(N = 3; Q = 2.328, p = 0.231) and within sound treat-
ments in Tank 2 vs. 1 (N = 3; Q = 2.430, p = 0.203).
Mean SGR was also higher in control tanks (0.87 ±
0.07) than in sound treatments (0.80 ± 0.07) but val-
ues did not differ significantly between treatments
(F1,3 = 4.724, p = 0.118).

3.6. Salmon blood analyses and histology

All blood and histological parameters from pool-
sampled animals (at experimental onset) were within
normal bounds for juvenile Atlantic salmon (Table 1).
At experimental completion, all blood and histo-
logical parameters remained within normal or ex-
pected values. Blood glucose and hematocrit levels
increased from the experimental outset to final sam-
pling in both treatment and controls. Mean blood
glucose concentration was higher in sound treat-
ments (112.8 ± 27.9 mg l−1) than in the controls (96.7
± 12.5 mg l−1); however, values were not significantly
different (F1,17 = 0.831, p = 0.413) with no tank effect
(Tank [Treatment] F4,17 = 2.825, p = 0.072). Lactate in
sound tanks (1.7 ± 0.5 mmol l−1) was lower than in
controls (2.0 ± 0.5 mmol l−1) but values did not differ
significantly (F1,15 = 0.144, p = 0.725) with no tank
effect (Tank [Treatment] F4,15 = 3.000; p = 0.072).
Treatment pH (7.2 ± 0.1) did not differ significantly
from controls (7.3 ± 0.1) (F1,17 = 0.333, p = 0.594) with
no tank effect (Tank [Treatment] F4,17 = 0.928, p =
0.479). Hematocrit in treatment fish (34.1 ± 3.3%) did
not differ significantly (F1,17 = 0.276, p = 0.626) from
controls (34.6 ± 4.2%) with no tank effect (Tank
[Treatment] F4,17 = 0.317, p = 0.860). Liver weight in
Treat fish (3.9 ± 0.4 g) did not differ significantly
(F1,17 = 0.186, p = 0.688) from controls (4.0 ± 0.2 g) with
no tank effect (Tank [Treatment] F4,17 = 0.352, p =
0.837). Hepatosomatic index in Treatment fish (1.5 ±
0.02%) did not differ significantly (F1,17 = 0.236, p =
0.652) from controls (1.4 ± 0.02 g) with no tank effect
(Tank [Treatment] F4,17 = 2.024, p = 0.154).

Plasma cortisol levels at experimental completion
were highly variable and higher in sound-exposure
treatments (34.7 ± 15.2 ng ml−1) than in controls
(30.44 ± 45.8 ng ml−1) but did not differ significantly
between treatments (F1,17 = 0.023, p = 0.886) with no
tank effect (Tank [Treatment] F4,17 = 2.108, p = 0.142;
Table 1).

4. DISCUSSION

High-intensity soundscapes created by aquacul-
ture production systems remain poorly understood.
There is a dearth of information on their effects on
growth performance, stress and immune status and
overall welfare of the high-value animals exposed to
them. The current study developed and applied novel
exposure experiments using playback of recordings
of commercial aquaculture production systems for
high-value global aquaculture species Atlantic salmon
Salmo salar and whiteleg shrimp Litopenaeus van-
namei. This study revealed no measurable impact on
cultured animals exposed to aquaculture system
soundscapes over a period of 8 wk. The measures
applied in the current study are broad and were cho-
sen to represent key performance, metabolic and
immunological parameters of interest for commercial
aquaculture production research. Herein, they are
discussed in relation to previous behavioural and
metabolic studies of sound impacts on fish and
invertebrates.

The sounds generated by both of these aquaculture
systems was relatively quiet (128 and 127 dB re 1 µPa)
when compared to other anthropogenic sources of
sound (Radford & Slater 2019). For example, commer-
cial ships have source levels ranging from 175 to
192 dB re 1 µPa2 in the frequency band 20 to 1000 Hz
(McKenna et al. 2013) and pile-driving activity gener-
ates sound exceeding 170 dB re 1 µPa (Würsig et al.
2000). The sounds generated by these 2 systems are
more consistent with recreational boat noise, which

| Physiological parameter | Lactate (mmol l−1) | pH | Liver weight (g) | Fish weight (g) | HSI (%) | Glucose (mg l−1) | Haematocrit (%) | Plasma cortisol (ng ml−1) |
|------------------------|-------------------|----|-----------------|----------------|---------|----------------|-----------------|-------------------------|
| Exp. onset              | 1.9 (0.9)         | 7.4 (0.1) | n.a.            | 179.8 (60.4)   | n.a.   | 62.8 (12.2)   | 16.6 (10.4)   | n.a.                    |
| Sound                  | 1.7 (0.5)         | 7.2 (0.1) | 3.9 (0.4)       | 262.0 (44.1)   | 1.5 (0.02) | 112.8 (27.9) | 34.1 (3.3)    | 34.7 (15.2)             |
| Control                | 2.0 (0.5)         | 7.3 (0.1) | 4.0 (0.2)       | 285.6 (34.8)   | 1.4 (0.02) | 96.7 (12.5)  | 34.6 (4.2)    | 30.44 (45.8)            |
ranges from 111 to 135 dB re 1 µPa depending on engine type and distance to the source (Nedelec et al. 2017, Mensinger et al. 2018). Chronic or acute sound exposure can result in different responses in terms of larval behaviour. For example, Nedelec et al. (2016) showed that the coral reef fish, Dascyllus trimaculatus, habituated to chronic exposure to recreation boat sound, but showed a significant stress response to multiple acute exposures to the same sound. Hence, the lack of stress responses observed both in whiteleg shrimp and Atlantic salmon in the present study maybe due to habituation to the sound playback.

Filiciotto et al. (2013) measured increased oxidative stress response in juvenile sea bream after 40 d exposure to onshore aquaculture sounds ranging up to 126 dB re 1 µPa and in control tanks with similar continuous sounds ranging up to 115 dB re 1 µPa. However, in the same study, reduced oxidative stress and higher growth performance was measured in animals exposed to highly variable soundscapes representing offshore aquaculture conditions and ranging up to 146 dB re 1 µPa (Filiciotto et al. 2013). Wysocki et al. (2006) were also able to detect significant differences in water cortisol concentration for European perch and common carp after short-term exposure to ship noise at 153 dB re 1 µPa. In the present study, 8 wk exposure to levels near 130 dB re 1 µPa did not cause significant increases in plasma cortisol in Atlantic salmon and did not significantly affect prophenoloxidase response in whiteleg shrimp. Behavioural and physiological hearing thresholds vary widely among species (e.g. Hawkins & Johnstone 1978, Amoser & Ladich 2005); however, the sound and playback levels in the current study are well within the hearing range of Atlantic salmon and that determined for a similar shrimp species, the common shrimp Palaemon serratus (Hawkins & Johnstone 1978, Amoser & Ladich 2005, Lovell et al. 2005, Ladich & Fay 2013). Hearing capacity in rapidly growing salmon may be reduced due to otolith malformation (Reimer et al. 2016, 2017). Nonetheless, it appears unlikely that the animals in the current study did not perceive the playback sounds at all. However, it is possible that the markedly (~14 dB re 1 µPa) lower and less variable (compared to previous experiments, e.g. Wysocki et al. [2006] used ship sounds ranging from 128 to 162 dB) SPLs in the current study did not elicit the strong behavioural or physiological responses observed in other studies (Wysocki et al. 2006). The SPLs used in the current study represent the prevailing acoustic habitat at aquaculture sites and the playbacks are actual recordings of those sites. These data may provide first reference values for maximum SPLs at sites and may assist in the development of guidelines to reduce stressful sound events.

It remains unclear whether Atlantic salmon and whiteleg shrimp habituated to the sound exposure in the present study of 56 d. A normalization of blood cortisol levels may have occurred as animals become habituated to higher sound levels. The current results suggest rather that the sound levels do not result in a cortisol response in the first instance. The equal growth, survival and metabolic responses indicate that the animals did not suffer significant or disadvantageous stress over the experimental period. In aquaculture systems, where exposure to soundscapes is a permanent and unavoidable factor for the fish, the longer-term response is highly applicable. Wysocki et al. (2006) suggested that high-level Gaussian or white noise at 153 dB re 1 µPa did not induce a stress response due to the lack of variability in noise level and type. Filiciotto et al. (2013) also used 5 alternating sounds at 5 varying SPLs to simulate offshore aquaculture noise and recorded reduced oxidative stress and increased growth in sea bream exposed for several months when sounds were more variable, despite being louder. The recordings applied in the current study are variable, particularly the salmon farm recordings. It is probable that the shrimp RAS system recording is not only quieter but also markedly less variable than those used in previous exposure studies (Radford & Slater 2019). However, this resembles the actually occurring sounds in German shrimp production systems.

High levels of sound exposure from pile-driving cause large-scale (hundreds to thousands of metres) avoidance responses in several salmonids and smaller scales (1 to 10 m) in shrimp, including whiteleg shrimp (Feist 1991, Zhou et al. 2016). In wild organisms, noise avoidance can markedly affect feeding, breeding or options for predator avoidance. While breeding and predators should play small roles in grow-out aquaculture, feeding disturbance due to sound would result in growth and survival impairment. In the current study, such impairment was not observed: growth rates were excellent for both species and did not differ between control and treatments. The current results for whiteleg shrimp strongly contradict Lagardère (1982), who reported that sound levels of 30 dB re 1 µBar/Hz at 25 to 400 Hz frequency range resulted in lower growth and reproduction rates and higher aggression levels with poorer feeding in tank-reared brown shrimp Crangon crangon. Apart from differences in claw count and the tail segment shape, brown shrimp do not
exhibit marked morphological differences to white-leg shrimp. However, the species is much smaller than white-leg shrimp (<30% of adult size) and exhibits different daytime behaviour in tanks with a distinct burying activity during high light periods (M. Slater pers. obs.). Responses of aquaculture species to sound, particularly variable sound, may be unexpected and complex, even positive in terms of feeding response. In a conditioning study, Meier & Horsemann (1977) were able to use daily 20 min buzzer sounds to induce a feeding response in blue tilapia Oreochromis aureus, which resulted in significant growth improvements over control fish. This result was obtained with robust, grow-out ready fish, as in the current study. Such robustness may not be given in earlier life stages, which may point to future areas of research.

Previous studies with larval fish indicate that sound exposure can affect growth, survival, reproductive success and even levels of aggression. Banner & Hyatt (1973) increased SPLs by 20 dB above controls and found reduced growth in sheepshead minnow Cyprinodon variegatus and longnose killifish Fundulus similis. In Atlantic cod Gadus morhua larvae, early hatched animals showed more rapid egg-sac depletion, increased startling events and reduced growth after 2 d of exposure to randomised and continuous sounds compared to a sound-free control (Nedelec et al. 2015). However, all growth and survival measures converged between treatment and controls after 16 d of exposure. The impact of sound on juvenile crustaceans has not been investigated, but the present results indicate this may be an area of future interest in terms of aquaculture soundscapes and limit-setting for future SPLs in hatcheries.

The current results indicate that actual soundscapes from sea pen and recirculating aquaculture facilities do not negatively affect growth, metabolism, health or immune response in the high-value species studied herein during the grow-out phase. Where soundscapes exceed the SPLs applied herein, the literature indicates that stress responses can be expected. The SPLs tested herein should act as threshold maximum SPLs and can be applied in a precautionary approach to ensuring soundscapes suitable for ensuring animal welfare is addressed in terms of sound in aquaculture. Further research regarding the impact on early juvenile animals particularly in tank/recirculating systems in hatcheries is recommended. Studies on sensitive adult stages such as broodstock at varying stages in the breeding cycle and on aquaculture species of potentially high sensitivity are recommended.

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