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Free-living *Neoparamoeba perurans* depth distribution is mostly uniform in salmon cages, but reshaped by stratification and potentially extreme fish crowding

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ABSTRACT: Understanding the spatio-temporal positioning of hosts and their parasites in freeliving states is useful in devising methods to diminish parasite encounters in animal production systems. We explored the potential for depth-based control methods of the amoebic gill disease (AGD) agent, Neoparamoeba perurans, in salmon mariculture systems by conducting: (1) depthstratified N. perurans water sampling surveys in 2 years, and (2) a mensurative experiment comparing depth distributions of *N. perurans* and salmon hosts in commercial salmon sea-cages. From water sampling mostly at marine salinities, N. perurans abundance (quantitative PCR-derived cells l⁻¹) varied among years, but overall, neither depth, time since freshwater bathing, temperature and salinity were predictors of *N. perurans* abundance. However, at 1 survey time, depth patterns in N. perurans abundance appeared during strong vertical salinity gradients following rainfall (at 1 site, salinity ranged between 14 and 35 g l^{-1}), with greater numbers of cells below a less saline surface layer. This suggested that salinity mediates N. perurans depth distribution during intermittent halocline development. Fish depth distribution monitoring revealed intense fish crowding, with local swimming densities up to 5 times stocking densities, typically at the surface at night. Simultaneously collected daytime water samples during low levels of fish crowding, with stock scattered amongst upper and lower cage sections, revealed no relationship between N. *perurans* and fish depth distributions. If intense fish crowding in narrow depth bands leads to high concentrations of N. perurans in cage environments and increased AGD risk, behavioural manipulations that vertically spread fish could be a successful AGD mitigation strategy.

KEY WORDS: Amoebic gill disease \cdot *Neoparamoeba perurans* \cdot *Salmo salar* \cdot Depth \cdot Swimming density \cdot Aquaculture

INTRODUCTION

The marine amoeba *Neoparamoeba perurans*, which causes amoebic gill disease (AGD), is one of the major parasites of concern in salmon sea-cage culture, with outbreaks recorded in all major salmonproducing regions (Oldham et al. 2016). In the most extreme case, summer–autumn AGD outbreaks have persisted in the Atlantic salmon *Salmo salar* industry within Tasmania since its inception 3 decades ago

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(Munday et al. 1990, 2001, Oldham et al. 2016). Despite the search for alternative controls and the start of programs to selectively breed salmon for AGD resistance (e.g. Kube et al. 2012), the same labour-intensive and partially-effective freshwater bathing treatments used to control AGD in the 1980s are still practiced today (Nowak 2012), with limited innovation of new control methods.

Many new preventive methods have surfaced for another important parasite, the salmon louse *Lepeophtheirus salmonis* (Dempster et al. 2009, Korsøen et al. 2012, Stien et al. 2012, 2016, Nilsen et al. 2017), after understanding the spatio-temporal depth distribution of its free-living infective larvae and salmon hosts in sea-cage environments (Heuch et al. 1995, Oppedal et al. 2011). Knowledge of how salmon hosts and free-living *N. perurans* distribute within seacages could therefore spur on the development of prophylactic techniques for *N. perurans*.

The depth-related swimming behaviour of salmon in sea-cages is well known in Norwegian coastal and fjord settings. Salmon adjust their vertical position by trading off between internal and external factors. Important environmental variables include light, temperature, salinity, dissolved oxygen, water currents and turbidity, while identified internal drivers consist of feeding or hunger status and the avoidance of perceived threats, such as predators (reviewed by Oppedal et al. 2011). In heterogeneous cage environments, salmon prefer to swim at specific depths, mostly driven by photo- and thermo-regulation (e.g. Oppedal et al. 2007). When fish feed and are hungry, they position themselves in the area that feed is dispensed, usually in surface waters (Oppedal et al. 2011). This information has been used to model and predict Atlantic salmon swimming depths and densities within cages in Norway (Føre et al. 2009, 2013). However, in SE Tasmania, Australia, caged Atlantic salmon swimming behaviour is yet to be examined, and differences may be expected due to disparate environmental conditions (e.g. strong tidal currents, warm, oxygen-deficient waters and higher light irradiance levels) and inner motivations of fish (e.g. seal presence, increased risk of oxygen and temperature stress and different feeding regimes).

By comparison, limited information exists on the spatio-temporal distribution of free-living *N. peru*rans. In the last decade, *N. perurans* was confirmed as the disease-causing agent of AGD (Young et al. 2007, 2008, 2014, Crosbie et al. 2012), a molecular method for detecting it in the marine environment was developed (Bridle et al. 2010, 2015, Wright et al. 2015b), and the first assessments of its free-living

distribution in salmon sea-cages were conducted (Bridle et al. 2010, Wright et al. 2015b, Hellebø et al. 2017). These assessments found that N. perurans occurs throughout the water column within farms holding AGD-affected stock in Australia (Bridle et al. 2010, Wright et al. 2015b) and in Norway (Hellebø et al. 2017). Further, free-living N. perurans are most abundant when clinical AGD levels are highest in stock and/or water temperatures are elevated (Wright et al. 2015b, Hellebø et al. 2017), and there have been occasions of higher abundance in the upper cage depths (Wright et al. 2015b, Hellebø et al. 2017). Surface concentrations of N. perurans have been hypothesised to result from either the swimming behaviour of caged salmon or the life history of this amoeba (Wright et al. 2015b). However, assessments have yet to compare depth-related free-living N. perurans concentrations with those of salmon in cages or take place during periods where strong vertical gradients in environmental variables exist that affect AGD risk, e.g. salinity and temperature (Clark & Nowak 1999, Adams & Nowak 2003, Oldham et al. 2016).

For parasites benefiting from host proximity for transmission, such as N. perurans (Nowak 2012), higher host density is theorised to increase the probability of a parasite encountering a new host. Positive relationships between host density and parasite intensity occur for this parasite type across a range of host taxa (Côté & Poulin 1995, Arneberg et al. 1998). This also appears to be the case for *N. perurans*, with AGD progressing faster at higher stocking densities for salmon held in tanks (Crosbie et al. 2010), and some evidence of this in sea-cages (Douglas-Helders et al. 2004). Given that caged salmon can crowd in narrow depth bands at many times the densities at which they are stocked (reviewed by Oppedal et al. 2011), local swimming densities may contribute to free-living *N. perurans* abundance and encounter risk in sea-cage environments. If tight-packing in narrow depth intervals is an important driver of N. perurans abundance, avoiding these situations through salmon behaviour or cage environment manipulations may improve AGD management, e.g. vertical light movements (Wright et al. 2015a).

Here, we surveyed the distribution of free-living *N. perurans* in commercial salmon cages of SE Tasmania using depth-dispersed water sampling and quantitative PCR (qPCR) methods in the AGD outbreak season (summer–autumn) of 2015 to add to data from the 2014 season (Wright et al. 2015b). This enabled a 2 yr data set to better decipher the relative importance of depth, as well as environmental and cage-specific variables in explaining *N. perurans* abundance in

salmon sea-cages. In addition, we investigated the depth-related behaviour of caged salmon and examined its influence on the depth distribution of free-living *N. perurans*. This involved a mensurative experiment using echo-sounders to continuously monitor the vertical distribution of caged salmon populations in concert with depth-stratified water sampling to quantify *N. perurans* using qPCR techniques.

MATERIALS AND METHODS

Neoparamoeba perurans depth survey

In the 2015 *N. perurans* depth survey, we surveyed *N. perurans* depth distribution in 9 commercial salmon cages (diameter = 38-54 m, circumference = 120-168 m, cage depth = 10-19 m) at 3 sites (3 cages site⁻¹) in the D'Entrecasteaux Channel, SE Tasmania, Australia (operated by TASSAL and Huon Aquaculture Company). Water samples were taken at depths of 0 (collected by slightly lifting the top of the Niskin bottle above the surface), 1, 2.5, 5 and 10 m in each cage at 4 times in 2015 on 20-22 January (time 1), 23-25 February (time 2), 18-26 March (time 3) and 29 April – 1 May (time 4) in 2015. We took single samples across the 5 depths, 9 cages and 4 times (for a total of 180 samples).

Fish vs. N. perurans depth experiment

In 2015, a fish vs. *N. perurans* depth experiment was conducted at a separate salmon farm location from *N. perurans* depth surveys. We measured the depth distribution of salmon and free-living *N. perurans* in 3 square commercial cages (square = 30×30 m, cage depth = 10 m) at the Northwest Bay lease (operated by TASSAL) in the northern end of the D'Entrecasteaux Channel, SE Tasmania, Australia. Fish depth was monitored for a ~24 h period from 12:00 to 15:00 h (GMT+10), towards the end of which water sampling for *N. perurans* abundance occurred at 0, 2.5, 5, 7.5 and 10 m depth in daylight hours on 18–19 February (time 1), 24–25 March (time 2) and 5–6 May (time 3). In total, 45 samples were taken (5 depths, 3 cages and 3 times).

N. perurans water sampling and quantification

For both the survey and experiment, water sampling for *N. perurans* was done as described by

Wright et al. (2015b). We positioned a 1.7 l Niskin bottle (General Oceanics) centrally at a quarter of the cage diameter within a cage above the transducer (fish and N. perurans depth experiment) or at the downstream end (survey of N. perurans depth). A rope system allowed the triggering of bottle closure at a range of depths by a cage-side operator (to avoid changes in normal fish behaviour). Collected water was poured from the sampler into 2×1 l sterilised plastic bottles, transported on ice and refrigerated. All water samples were filtered within 24 h using filters (1.2 µm pore size, 47 mm diameter, GF/C Whatman glass microfibre, Sigma-Aldrich), a filter holder/receiver (Nalgene™, Thermoscientific) and a Rocker 300 vacuum pump (JAVAC) operated at 30 kPa. Filters were inserted into 5 ml vials with 1 ml lysis buffer (4 M Urea, 1% SDS, 0.2 M NaCl, 1 mM sodium citrate) and frozen at -20°C. Real-time gPCR assays for N. perurans were then performed to calculate N. perurans cell abundance from thawed samples via a CFX Connect PCR Detection System (Bio-Rad). This distinguished a N. perurans-specific 18S rRNA gene sequence and estimated the number of cells based on the conversion: 2880 copies = 1 N. perurans cell (Wright et al. 2015b). N. perurans cell abundance was recalculated to cells l^{-1} for each sample to account for differences in sample volumes.

Environmental and AGD-related variables

At the time of water sampling at a cage in both the survey and experiment, salinity and temperature were recorded using a CTD profiler (model SD204, SAIV-AS) lowered through the water column from 0 to 10.25 m. Recordings were then averaged at 0.5 m intervals. We also gathered information on days since the last freshwater bath, stocking density and AGD prevalence (percentage of 30-40 fish with macroscopic AGD spots or lesions). We reported on pre-bath AGD prevalence, which was measured before the most recent freshwater bath in a given cage (Wright et al. 2015b). Historical environmental data were obtained to help explain inter-annual patterns in N. perurans abundance. This consisted of daily sea surface temperature measurements at the mouth of the Huon Estuary (data provided by Huon Aquaculture Company, only available from March 2014) and daily rainfall recording at Dover (Australian Bureau of Meteorology weather station number 094020).

Fish depth monitoring

In the fish vs. N. perurans depth experiment, salmon depth was measured and logged via an echo-integration system (model CageEye mk.II, Lindem Data Acquisition) connected to 3 transducers (46° beam, 50 kHz, 0.001 s pulse, 1 s echo listening, 3 s pulse interval) positioned ~7 m underneath the bottom of each study cage (Bjordal et al. 1993). The transducers pointed up centrally at a quarter-distance spacing (7.5 m of 30 m wide cages) from a cage edge running parallel to the tidal current direction. This was expected to capture a representative sample of the depth distribution for schooling fish inside large commercial cages, and capture fish depth patterns at the position of water sampling for N. perurans. Echo intensity (EI) recordings were condensed to average values every minute, and from 0.07 m depth increments between 0.25 and 10 m to average values every 0.5 m. Across the 20 depth intervals (n), data were then converted to relative echo intensity (ER) using the formula

$$ER_{n} = EI_{n}^{-(0.01 \times \Sigma EI_{1-20})}$$
(1)

From this, observed fish densities (OFD or local swimming density, kg $\rm m^{-3})$ at each depth interval were calculated by

$$OFD_n = B \times ER_n \times V_n^{-1}$$
(2)

where B is the total cage biomass, and V is the cage volume within a 0.5 m band. Values were allocated to day or night categories according to local sunset and sunrise times of 20:15 and 6:36, 19:17 and 7:19, and 17:09 and 7:07 h for times 1, 2 and 3, respectively. Average minute data were used (see Figs. 3 & 5); however, day and night OFD data were further condensed to average hourly values in preparation for statistical analyses of maximum OFD (OFD_{max} or local swimming density at the most preferred swimming depth), the depth of OFD_{max} and the preference index (PI) or the guantified magnitude of preference for specific depths (Oppedal et al. 2007). For each hour, the PI was obtained from the equation

$$PI = \sum AER_n \div \sum D_n \tag{3}$$

for which $\sum AER_n$ represents the sum of residual echo intensities above what was expected under a uniform fish distribution (1/20 depth intervals = 0.05) and $\sum D_n$ is the count of those depths (Oppedal et al. 2007).

Statistical analyses

The importance of predictor variables in explaining N. perurans cell abundance within salmon cages at survey times in 2015 were examined using generalised additive mixed model (GAMM) analyses (Lin & Zhang 1999) in R (R Core Development Team, version 3.1.0) via the mgcv package. GAMMs are nonparametric and use a smooth function to estimate predictor effects, in contrast to generalised linear mixed models (GLMMs) that fit a linear function and rely on normally distributed data. We opted for GAMMs in favour of GLMMs used in previous work (Wright et al. 2015b), because 2015 data were nonnormal even after data transformation as a result of low N. perurans cell abundance. Untransformed N. perurans cell abundance (cells l^{-1}) was set as the response variable, and predictors used as fixed effects were depth, time since freshwater bathing, water temperature and salinity (Wright et al. 2015b). Site was included as a random factor. Model selection was determined from Akaike's information criterion (AIC) (Akaike 1973). The best model was one with the lowest AIC value, and a simpler model was considered better than one with an additional variable if the difference in AIC was ≤ 2 (Burnham & Andersen 2002). We also used Akaike weights (w_i) to evaluate the relative likelihood of each model, with higher weights indicating improved support (Burnham & Andersen 2002). These analyses were repeated for a 2014 dataset pooled across 3 times in March, April and June from a previous survey study (Wright et al. 2015b), a 2015 dataset pooled across 4 times in January, February, March and April-May from this survey study and finally a multi-year survey dataset pooled across all 7 times. We also assessed inter-annual differences in N. perurans cell abundance between 2014 and 2015 surveys using a Mann-Whitney U-test.

For time 3 of the fish vs. *N. perurans* depth experiment (the only time *N. perurans* cells were detected), we examined differences in $\log_{10}(x + 1)$ -transformed OFD measurements at the depth of *N. perurans* water samples from all replicate cages between those that were positive and negative using *t*-tests. These were performed for the OFD recorded at the exact minute of sampling (0 min), the previous 1 min, or average OFD values from the previous 10, 60 (1 h), 360 (6 h) or 1440 min (1 d) of sampling within a given cage. OFD at the sampling depth in a cage was calculated by averaging OFD recordings in the 1.25 m above and below the sampling depth (0.25–1.25 for 0 m, 1.25–3.75 for 2.5 m, 3.75–6.25 for 5 m, 6.25–8.75 for 7.5 m and

8.75–10.0 m for 10 m). Mean hourly OFD_{max} values with a $log_{10}(x + 1)$ transformation pooled from replicate cages were compared between time of day (day or night) and monitoring period (time 1–3) using a 2-factor ANOVA. Significant results were followed by Tukey HSD post hoc comparisons to determine statistically different groups. Normality and equality of variance was checked using Shapiro-Wilk tests and residual plots, respectively, leading to data transformations where necessary. Significance was set at $\alpha = 0.05$, and error values refer to standard errors.

RESULTS

Neoparamoeba perurans depth survey

Preceding heavy rainfall caused a less saline (minimum of 14 g l⁻¹) and warmer surface layer (maximum) of 20°C) in study cages at time 1. Less pronounced stratification was present at time 2 (minimum salinity of 29 g l^{-1}) and became mostly absent thereafter (Fig. 1). Mean abundance (in cells l^{-1}) of *N. perurans* cells was low at times 1 (0.04 \pm 0.02), 2 (0.10 \pm 0.07), and 3 (0.02 \pm 0.02), but increased at time 4 (0.35 \pm 0.24; Fig. 1). The comparatively elevated N. perurans cell abundance at time 4 was matched by the highest bathing frequency (Table 1). Based on pre-bath gill scores, AGD was less prevalent at time 4, although these scores were potentially misleading because of changing bathing frequency and environmental conditions throughout the survey (Table 1, Fig. 1). Small progressive increases in stocking densities were seen in study cages over time (Table 1).

At time 1 when brackish water was present at the surface, GAMM analyses revealed that the model using depth best explained *N. perurans* cell abundance (Table S1, in the Supplement at www.int-res. com/articles/suppl/q009p269_supp.pdf). The model coefficient indicated that *N. perurans* cells were more abundant deeper in the water column (df = 6.4, $R^2 = 0.10$, p = 0.02; see Table S1 in the Supplement. In contrast, at times 2–4 when vertical temperature and salinity gradients weakened or vanished, no models performed better than the null model in explaining *N. perurans* abundance (Tables S2–S4 in the Supplement).

Inter-annual variation in N. perurans depth

N. perurans cells were more abundant in 2014 $(1.87 \pm 0.66 \text{ cells } l^{-1})$ than in 2015 $(0.13 \pm 0.06 \text{ cells } l^{-1};$

W = 16696, p < 0.0001; Fig. 2). This was in conjunction with substantially less rainfall in 2014 (271 mm from January-June) relative to 2015 (507 mm from January-June; Fig. 2). Depth patterns also varied between years, with greater numbers of N. perurans cells in surface waters at time 1 in 2014 (after minimal rainfall) and the opposite at time 1 in 2015 (after 116 mm of rainfall on 14-15 January 2015; Fig. 2). However, at all other times, no depth-related pattern existed. GAMM analysis revealed that temperature weakly explained N. perurans cell abundance in 2014 (df = 6.8, R^2 = 0.04, p = 0.01; Table S5 in the Supplement), with more cells at higher temperatures (Fig. 2). However, for 2015 and the multi-year dataset (2014 and 2015 combined), the addition of predictors did not help explain N. perurans cell abundance beyond null models (Tables S6 & S7 in the Supplement).



Fig. 1. Mean (± SE) of temperature and salinity at 0.5 m depth increments (solid circles) and *Neoparamoeba perurans* cell abundance at 0, 1, 2.5, 5 and 10 m (open circles represent individual data points) in surveyed Atlantic salmon *Salmo salar* cages at times 1–4 in 2015 (1: 20–22 January; 2: 23–25 February; 3: 18–26 March; 4: 29 April–1 May)

Table 1. Mean $(\pm SE)$ of pre-bath amoebic gill disease (AGD) level, time since freshwater bathing and stocking density of Atlantic salmon *Salmo salar* for surveyed salmon cages at times 1–4 (see Fig. 1 for exact times). Ranges are displayed in parentheses; all values are based on 9 cages at each time point

Time	Pre-bath AGD	Time since	Stocking density
	prevalence (%)	bathing (d)	(kg m ⁻³)
1	$50.7 \pm 8.5 (23-100)$	$27.0 \pm 4.3 (9-47)$	$5.3 \pm 0.5 (2.6-6.9)$
2	$54.1 \pm 6.8 (15-100)$	$28.8 \pm 4.4 (6-48)$	$5.6 \pm 0.5 (3.1-7.6)$
3	$60.6 \pm 4.5 (30-87)$	$25.9 \pm 3.6 (14-49)$	$5.7 \pm 0.5 (2.8-7.5)$
4	$38.9 \pm 2.6 (20-72)$	$17.8 \pm 3.2 (7-35)$	$6.6 \pm 0.6 (3.8-8.3)$

Fish vs. N. perurans depth experiment

Fish often displayed a bimodal depth distribution with peak local swimming densities (OFD_{max}) in upper (0–4 m) and lower (6–10 m) sub-groups (Figs. 3 & 4). At time 1, when temperatures were >18°C at 0–4 m and ~1°C cooler at 6–10 m, fish positioned in both the upper and lower sub-groups throughout day and night periods, with increased selection of deeper water during daylight hours (Figs. 3 & 4). At time 2, in the absence of thermal stratification in waters >16°C, fish concentrated predominantly in the upper 4 m regardless of light conditions (Figs. 3 & 4). At time 3, also without vertical temperature differences at ~13°C, fish were densely packed in the upper cage depths at night and then split between the upper and lower sub-groups in daylight (Figs. 3 & 4).

Mean OFD_{max} values varied with light conditions $(F_{1,228} = 103.3, p < 0.0001)$ and over time $(F_{2,228} = 86.9, p < 0.0001)$, with interactions $(F_{2,228} = 12.6, p < 0.0001)$. Higher OFD_{max} recordings occurred at night relative to daylight hours, excluding time 1 (p < 0.05; Table 2). OFD_{max} levels were greater on Nights 2 and



3 than in all other periods, values on Days 2 and 3 were higher than those on Day 1, and recordings on Day 2 were above those on Night 1 (p < 0.05; Table 2). Similarly, mean preference indexes were elevated (>2), indicating increased crowding at specific cage depths, during Nights 2 and 3 and on Day 2 (Table 2). The highest OFD_{max} recorded using hourly average data was 52 kg m⁻³ or 5.2 times the mean stocking density on Night 3 (Table 2). Based on minute average data, maximum OFD values reached 116 kg

 m^{-3} or 14.2 times the mean stocking density on Night 2 (Fig. 4; Table 2).

N. perurans cells were undetected in depth-stratified water samples at times 1 or 2, and were only found in low abundance at time 3 (Fig. 4). Time 3 coincided with homogeneous salinity and temperature conditions throughout cage depths and low levels of fish crowding (Fig. 4). Comparisons of mean OFD measurements at the sampling depth between samples with and without N. perurans cells at time 3 were not statistically different (Fig. 5). This was the case for mean OFD recordings at 0 min of samples being taken when the largest separation occurred (present vs. absent: 11.5 and 9.8 kg m⁻³, t = -1.6, p = 0.1), and recordings in the 1, 10, 60, 360 (6 h) or 1440 min (1 d) prior to sampling (t = -0.05 to -1.1, p > 0.3; Fig. 5). Of note, a 7.5 m water sample, corresponding to the highest mean OFD at sample depth at 0 min (25.3 kg m⁻³ or 2.6 times the stocking density) and in the previous 10 min (15.8 kg m^{-3} or 1.6 times the stocking density), contained the highest N. perurans cell abundance recorded (2.0 cells l⁻¹ compared to < 0.7 cells l^{-1} in other samples).

> Fig. 2. Mean (± SE) Neoparamoeba perurans cell abundance (filled circles) at each time in surveyed Atlantic salmon Salmo salar cages in the 2014 and 2015 surveys. When depth was the main explanatory factor in N. perurans cell abundance (determined by GLMM or GAMM analyses), the direction of depth effects is displayed (up arrow for more N. perurans cells in surface waters, and down arrow for greater numbers at lower cage depths). Daily rainfall (Dover station 094020, bars) and water temperature at 0 m (Roaring Beach lease provided by Huon Aquaculture Company, lines) are plotted from January-July in both years. No temperature data were available in January and February 2014

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Fig. 3. Observed fish density (OFD; colour coded) every minute between 0.25 and 10 m depth over ~24 h periods in each of 3 Atlantic salmon Salmo salar cages at times 1–3 in 2015 (1: 18–19 February; 2: 24–25 March; 3: 5–6 May. The panel above each graph represents night- (black bars) and daytime (open bars) based on local sunrise and sunset times, and the boxes within the graphs indicate periods during which water sampling for Neoparamoeba perurans took place

DISCUSSION

Depth patterns in *Neoparamoeba perurans* abundance were mostly absent within salmon cages. In the 2014 season after low rainfall, surface concentrations of *N. perurans* were seen at 1 out of 3 survey times (Wright et al. 2015b). In the 2015 season after high rainfall, *N. perurans* were deeper in the water column during strong vertical gradients in salinity and temperature, but with uniform distribution at the remaining 3 survey times. GAMM analyses for surveys from each year and both years combined confirmed that

depth was not an important driver of amoebae abundance. Caged salmon usually displayed their typical diurnal behavioural pattern (Oppedal et al. 2011) of deep daytime and shallow nighttime swimming, although they also occupied surface waters throughout the day (possibly explained by hunger levels) and lower cage depths at night (if these were of preferred temperature). In this study, local swimming densities of caged salmon were unrelated to *N. perurans* presence; further work is needed to test if this remains the case when *N. perurans* abundance and salmon swimming densities are high.



Fig. 4. Vertical patterns in environmental conditions (salinity and temperature as solid circles, mean (± SE) at 0.5 m intervals), observed fish density (OFD; percentage of maximum OFD at each 0.5 m interval) during day- (open bars) and nighttime (solid bars), and *Neoparamoeba perurans* cell abundance (open circles) at times 1–3 in 2015 (see Fig. 3 for exact times)

Surveys in 2015 for *N. perurans* in salmon seacages did not detect high cell abundance in surface waters reported in 2014 at the same locations (Wright et al. 2015b). Furthermore, when data from 2014 and 2015 were scrutinised together, depth-related patterns of *N. perurans* in the water column were infrequent at specific survey times. In 2015, more cells were in deeper cage depths at 1 survey time, which was associated with brackish surface water conditions following heavy rainfall (down to 14 g l⁻¹ at 0 m). This trend was also reported in an earlier assessment of the depth distribution of *Paramoeba* and *Neoparamoeba* spp. in AGD-affected salmon cages at a location with a halocline present and salinities of 28 g l⁻¹ at 0 m (Douglas-Helders et al. 2003). Freshwater runoff exiting the river mouth after rainfall may phys-

Table 2. Mean (± SE) hourly maximum observed fish density (OFD_{max}) values and preference indexes in triplicate Atlantic salmon Salmo salar cages during the day and at night for each of the 3 times (see Fig. 3 for exact times). Mean (± SE) stocking densities at each time are also displayed. Ranges are provided in parentheses. Letters for OFD_{max} values indicate statistically different groups revealed from Tukey HSD comparisons following a 2-factor ANOVA using time and light conditions as factors, and OFD_{max} as the response variable

Time	OFD _{max} (kg m ⁻³) Night	Preferer	ice index ——— ——— Night ————	Stocking density (kg m ⁻³)
1 2 3	$\begin{array}{l} 13.0 \pm 0.4^{a} \ (9.0{-}21.5) \\ 19.6 \pm 1.1^{c} \ (7.6{-}32.5) \\ 16.1 \pm 0.3^{bc} \ (13.1{-}19.7) \end{array}$	$\begin{array}{l} 15.4 \pm 0.7^{\rm ab} \ (9.1{-}22.6) \\ 26.8 \pm 1.4^{\rm d} \ (10.2{-}47.3) \\ 29.2 \pm 7.1^{\rm d} \ (18.9{-}52.0) \end{array}$	$\begin{array}{c} 1.15 \pm 0.11 \; (0.12 - 4.26) \\ 4.41 \pm 0.23 \; (0.76 - 7.48) \\ 1.48 \pm 0.06 \; (0.76 - 2.01) \end{array}$	$\begin{array}{l} 1.76 \pm 0.70 \; (0.40 - 2.95) \\ 6.18 \pm 0.31 \; (0.75 - 9.97) \\ 4.91 \pm 0.25 \; (2.10 - 10.42) \end{array}$	$\begin{array}{c} 8.5 \pm 0.6 \; (7.5 - 9.4) \\ 8.3 \pm 0.5 \; (7.5 - 9.1) \\ 10.2 \pm 0.2 \; (9.9 - 10.5) \end{array}$



Fig. 5. Mean (± SE) of observed fish density (OFD) values calculated at the minute of sampling or from the previous 1, 10, 60 (1 h), 360 (6 h) or 1440 min (1 day, cells from left to right) within the 1.25 m above and below water samples where Neoparamoeba perurans was absent (open bar) or present (grey bar) at time 3 (5–6 May 2015)

ically push *N. perurans* out the surface brackish layer. Concomitantly, free-living *N. perurans* using contractile vacuoles to maintain inner cell osmolality equivalent to full-strength seawater (Lima et al. 2016) would become denser than surrounding water at lower salinities, potentially causing them to sink out of such layers. Buoyancy changes could allow marine amoebae to move away from unfavourable low-salinity surface conditions. Overall, strong vertical gradients in salinity were only temporary and it was most common that *N. perurans* distribution did not show distinct depth patterns (Wright et al. 2015b).

Fewer free-living N. perurans were observed in the months after a major rainfall event in 2015. That year, rainfall appeared to overrule the influence of temperature on N. perurans abundance detected in 2014 (Wright et al. 2015b). AGD outbreak magnitude is reduced or avoided under brackish surface water conditions (Clark & Nowak 1999, Adams & Nowak 2003, Oldham et al. 2016). This likely eventuates for 2 reasons. First, as discussed above, increased river discharge may flush N. perurans out of estuaries particularly at upper cage depths, reducing free-living N. perurans abundance and new AGD infections. Second, gill-attached N. perurans could dislodge or be adversely affected when hosts briefly or persistently reside in brackish surface water following rainfall. Increased shedding of mucus from the gills of salmon occurs during prolonged reductions in salinity (Roberts & Powell 2003, 2005), and may be an important mechanism in N. perurans detachment. N. perurans attached to plastic substrates subjected to low salinities (28 g l^{-1}) either remain there using contractile vacuoles for osmotic regulation (Lima et al. 2016), or detach and assume floating rounded forms that are not expected to reproduce (Martin 1985, Cowie & Hannah 2006). Extended low-salinity exposures will eventually cause lysis of N. perurans (Powell & Clark 2003). Salinity reductions appear to diminish the abundance of free-living *N. perurans* in cage environments and control AGD outbreaks.

In SE Tasmania, the depth-related behaviour of caged salmon generally resembled that found in previous assessments in Norway (reviewed by Oppedal et al. 2011). Fish tended to occupy shallow depths at night and moved deeper during the day, which is diel behaviour widely reported in caged salmon (see Oppedal et al. 2011). However, during periods of thermal stratification, fish chose to swim at depths closest to their preferred optimal temperature range of 16-18°C (Johansson et al. 2006, 2009) for thermoregulation and intrinsic physiological performance benefits, regardless of light conditions (Oppedal et al. 2007, Stien et al. 2014). At one time, fish resided only near the surface throughout the day and night. This was unlikely motivated by thermal conditions, because vertical temperature differences were much less than those shown to drive depth selection of 0.2°C (Oppedal et al. 2007), and, at 16.2°C, these temperatures fell within the optimal range for Atlantic salmon. Instead, fish may have been underfed and hungry, occupying upper cage depths in anticipation of surface-delivered feed (Oppedal et al. 2011). Reduced feeding by farmers during high water temperatures and low oxygen levels, to minimise fish metabolic rates and oxygen consumption (Forsberg 1997), might have occurred prior to fish depth monitoring and explained surface-dwelling fish. Excluding this observation of daytime crowding at the surface, the tightest packing of caged salmon was at night, also in shallow waters.

Observed fish density effects on free-living *N. perurans* presence were not detected in our mensurative experiment in commercial salmon sea-cages. However, very few *N. perurans* were detected, suggesting they were shed from gills infrequently and few were present in the water column. Further, at the only time salmon swimming densities and *N. perurans* abundance could be compared, salmon were evenly partitioned between the upper and lower cage subgroups and were not crowding in any depth band. Consequently, our findings cannot be easily extrapolated to

conditions at the height of AGD outbreaks during intense fish crowding and high N. perurans abundance in the water column. The depth distribution of fish and free-living N. perurans in SE Tasmania indicate that a link is possible at those times. In homogeneous cage environments of ~16.3°C in 2014, free-living N. perurans were abundant in surface waters (Wright et al. 2015b). Analogously, during 16.2°C throughout depths in 2015, fish positioned in the upper cage depths during the most intense day-time crowding (mean packing index of 4.4, compared to <1.5 at other times). Hence, fish may have swum at the surface and driven the high *N. perurans* numbers there under identical conditions in the previous year. Future investigations examining the connection between fish and N. perurans depth distributions in cage environments should focus on the peaks of AGD outbreak (during low rainfall and high water temperatures) and extremes in fish crowding (mostly during night hours at the surface in this study).

CONCLUSIONS

From a multi-year dataset, we revealed that freeliving Neoparamoeba perurans are mostly ubiquitous throughout depths in salmon sea-cages and more abundant at high water temperatures when preceded by low rainfall. This implies that AGD cannot be avoided by excluding salmon from specific cage depth bands using behavioural and environmental manipulations, as is the case for *Lepeophtheirus* salmonis salmon lice (Dempster et al. 2009, Korsøen et al. 2012, Stien et al. 2012, 2016, Nilsen et al. 2017). However, our results do not preclude the possibility that behavioural and environmental approaches could be applied to reduce extreme local fish swimming densities, which were most common at night. Strategic deployment of depth-dispersed static and vertically moving artificial night lighting arrangements, that evenly spreads fish throughout cage depths, could slow AGD progression during outbreaks and improve salmon welfare more broadly (Juell et al. 2003, Oppedal et al. 2011, Wright et al. 2015a). While lights placed at single, static depths are in standard use to control maturation and the modifications we suggest to light placement, not intensity, are unlikely to alter stress levels in caged salmon, the repercussions of any modification to cage environments or salmon behaviour require careful assessment. Our study highlights the importance of understanding how hosts and parasites distribute spatio-temporally in farm environments to discover new parasite management tools.

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