1. INTRODUCTION

Gill health is fundamental to the success of finfish aquaculture. These delicate organs are responsible for gas exchange, acid–base balance, waste excretion and osmoregulation, yet are constantly exposed to a variety of microorganisms and environmental insults which can result in various gill diseases (Evans 2005, Mitchell & Rodger 2011). While the aetiologies of many disease pathologies are attributed to single pathological agents, the drivers of gill disease can be more complex (Gjessing et al. 2017, Herrero et al. 2018). A multifactorial approach to disease occurrence that considers the impact of environmental factors and microbial community structure, rather than focusing on a single agent, may therefore be a ra-
tional approach to gain a better understanding of disease in marine ecosystems (Egan & Gardiner 2016) and, in this case, gill disease.

Amoebic gill disease (AGD) is a globally significant disease in marine aquaculture and remains the main health issue challenging Atlantic salmon Salmo salar aquaculture in Tasmania, Australia. AGD is caused by Neoparamoeba perurans (syn. Paramoeba perurans; Feehan et al. 2013), a free-living protozoan ectoparasite which colonises the gills of Atlantic salmon in the marine phase of production (Young et al. 2007, Crossbie et al. 2012). Attachment of N. perurans causes lamellar fusion, epithelial hyperplasia, oedema and ultimately mortality when left untreated (Adams & Nowak 2001). While it is understood that salinity, water temperature and stocking density play a role in AGD prevalence and severity (Oldham et al. 2016), all the drivers of infestation and correlates of the associated gill pathology are not yet understood. This could contribute to inefficiencies in disease prediction and control during Atlantic salmon marine grow-out.

In addition to N. perurans, a diversity of other amoeba species colonise the gills of AGD-affected Atlantic salmon, with up to 11 other species having been isolated from farmed salmon in Tasmania, including the species of the genera Paramoeba, Vexillifer, Pseudoparamoeba, Vannella and Nolandella (Howard 2001, English et al. 2019). However, it is not understood whether these accompanying amoeba species are parasitic or commensal or if changes to the Amoebozoan community structure influence disease onset or progression. Moreover, the seasonal abundance of these amoebae and whether their presence correlates with AGD prevalence and severity of pathology is also unknown. Previous studies of gill-associated Amoebozoa identified the amoebae after isolating and culturing in vitro (Howard 2001, Birmingham & Mulcahy 2007, English et al. 2019). These methods likely generate a skewed assessment of infection prevalence and abundance because they are biased towards amoebae that are more capable of growing in the chosen culture conditions. A more accurate, sensitive and high-throughput technique for detecting and quantifying particular microorganisms is quantitative PCR (qPCR) (Purcell et al. 2011). There are 3 published qPCR assays for species-specific detection of N. perurans (Bridle et al. 2010, Fringuelli et al. 2012, Downes et al. 2015). These assays are versatile tools for research, being used to answer a variety of AGD-related questions, including retrospective validation that N. perurans was the main causative agent in the earliest AGD outbreaks in Ireland (Downes et al. 2018) and confirmation that Atlantic salmon can be experimentally infected from lumpfish Cyclopterus lumpus, a cleaner fish which is often cohabited with farmed salmon to control sea lice (Haugland et al. 2017). Despite these assays being highly informative and adaptable tools for disease and ecological research, there are no qPCR assays designed for the variety of other marine amoebae which colonise AGD-affected fish.

In a recent study, we described 18S rRNA sequences from many of the amoebae which colonise Atlantic salmon gills in Tasmania (English et al. 2019). These sequences are ideal for developing molecular detection assays which can be used to explore the potential role the Amoebozoan community plays in AGD onset and severity. Here, we design and validate 5 new amoeba qPCR assays using these 18S rRNA sequences. These assays, along with an existing N. perurans assay (Downes et al. 2015), were then used to assess amoeba prevalence and abundance on the gills of farmed Atlantic salmon over a 1 yr period from 2 different farm sites. This longitudinal survey aimed to identify relationships between specific amoeba taxa and farm sites, time points and gross gill pathology.

2. MATERIALS AND METHODS

2.1. Gill swab sample collection

Every 3 mo, from May 2017 to February 2018, during routine commercial gill checks at Tassal Operations Tasman and Killala leases (Fig. 1), large groups of Atlantic salmon Salmo salar were seined and crowded to ensure the fish were haphazardly mixed, then 40 fish were sampled from the crowd. The 2 farm sites have contrasting environmental conditions, with the Tasman lease being more oceanic, located closer to open water, and the Killala lease more brackish, located in the lower Huon Estuary. The average monthly water temperature and salinity at 5 m before each sampling event was obtained from Tassal Operations.

During sampling, fish were anaesthetised with 15 ppm of AQUI-S, then their gross gill pathology was scored according to Taylor et al. (2009) on a scale of zero (no gross lesions) to 5 (lesions covering >50% of gill surface). All 16 gill surfaces were then swabbed (Westlabs) in situ, with one swab used for the left gill arches and one for the right arches to maximize the DNA yield. The left and right swabs for each fish were pooled and stored in RNALater (Ambion) at room temperature for a minimum of 2 d and then at −20°C until processed for DNA extrac-
tion. Other information collected included months since the fish were transferred to sea, the average fish weight and the number of days since the fish were last treated for AGD (2 to 3 h freshwater bath). The case history of each sampling event is summarised in Table 1. All sampling events occurred as planned apart from February 2018 at Killala as the site was de-stocked as part of standard commercial operations. Sampling of 280 salmon in this study was approved by the CSIRO Queensland Animal Ethics Committee (AEC number 2017-16).

### 2.2. DNA extraction from gill swabs

DNA was extracted from the preserved gill swab samples using a plate-based DNeasy Blood and Tissue Kit (QIAGEN) according to the manufacturer’s instructions. Extracted DNA was quantified with a Nanodrop ND-1000 spectrophotometer (Life Technologies), then a working stock (30 ng µl⁻¹) of each sample was prepared using an epMotion 5070 liquid-handling robot (Eppendorf) and stored at −20°C.

### 2.3. TaqMan qRT-PCR assay design

Five TaqMan MGB probes and PCR primer sets were designed based on the 18S rRNA gene sequences of amoeboae previously isolated from AGD-affected Atlantic salmon (English et al. 2019). The chosen amoebae for assay design included Nolanella sp. (strain MX5, accession number MH535951), Pseudoparamoeba sp. (MX1, MH535967), Paramoeba eilhardi (106KRT, MH535952), Vannellida C2 (149SVA, MH535956) and Vannellida C3 (136SVA, MH535954). From here on, these assays, in the order listed above, will be referred to as Nol, Pse, ParE, VanC2 and VanC3. Based on an alignment of 18S rRNA gene sequences from these species and Neo-paramoeba perurans (MH535959), the primer and probe sets were designed on Primer Express 3.0.1 (Life Technologies) according to the guidelines. Before ordering the probes, the amplicon derived from PCR using each primer set was sequenced according to English et al. (2019) to confirm the correct sequence was being amplified. Primers were ordered from GeneWorks and probes from Thermo Fisher Scientific.

<table>
<thead>
<tr>
<th>Farm site</th>
<th>Sample time</th>
<th>Months since ocean transfer</th>
<th>Average fish weight (g)</th>
<th>Days since last freshwater bath</th>
<th>Water temp at 5 m (°C)</th>
<th>Salinity at 5 m (ppt)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Killala</td>
<td>May 2017</td>
<td>1</td>
<td>225</td>
<td>no bath</td>
<td>13.6</td>
<td>31.7</td>
</tr>
<tr>
<td></td>
<td>Aug 2017</td>
<td>3</td>
<td>719</td>
<td>25</td>
<td>11.9</td>
<td>31.9</td>
</tr>
<tr>
<td></td>
<td>Nov 2017</td>
<td>7</td>
<td>2530</td>
<td>64</td>
<td>14.7</td>
<td>31.1</td>
</tr>
<tr>
<td></td>
<td>Feb 2018</td>
<td>NA</td>
<td>NA</td>
<td>NA</td>
<td>NA</td>
<td>NA</td>
</tr>
<tr>
<td>Tasman</td>
<td>May 2017</td>
<td>11</td>
<td>2365</td>
<td>48</td>
<td>12.1</td>
<td>34.5</td>
</tr>
<tr>
<td></td>
<td>Aug 2017</td>
<td>3</td>
<td>639</td>
<td>35</td>
<td>10.2</td>
<td>31.1</td>
</tr>
<tr>
<td></td>
<td>Nov 2017</td>
<td>7</td>
<td>1618</td>
<td>37</td>
<td>15.5</td>
<td>32.0</td>
</tr>
<tr>
<td></td>
<td>Feb 2018</td>
<td>10</td>
<td>2734</td>
<td>50</td>
<td>17.5</td>
<td>31.8</td>
</tr>
</tbody>
</table>
Scientific. The optimum probe and primer concentration for a single-plex assay was then determined with a series of experiments following the Thermo Fisher Scientific standard protocols using amplicon-specific plasmid DNA as template (Thermo Fisher 2010). The amplicon-specific plasmid DNA was generated as described previously (English et al. 2019).

2.4. Validation of reaction efficiency, sensitivity, specificity and reproducibility

Assay validation was performed according to Downes et al. (2015) and Bustin et al. (2009). The following procedures were conducted separately on each of the 5 qPCR assays. To determine the efficiency of the assays, amplicon-specific plasmid DNA was serially diluted 10-fold and amplified in quadruplicate qPCR reactions. Only dilutions that provided a cycle threshold (Ct) value in all replicates were used to generate a standard curve. Amplification efficiency was calculated based on the Ct slope method (Efficiency = \[10^{(−1/slope)} − 1\]), and the linearity was demonstrated with the coefficient of determination (R²). The equivalent number of 18S rRNA copies corresponding to each standard curve was then determined by measuring the concentration of the plasmid DNA with a NanodropND-1000 spectrophotometer (Life Technologies) and then submitting this value and the length of the amplicon insert and vector to the DNA copy number calculator (Staroscik 2004). The 10-fold dilution series was then used to determine the limit of detection (LOD) for each assay. For this purpose, the lowest dilution that provided a Ct value in all replicates underwent a 2-fold dilution and was tested in quadruplicate. The mean of the lowest dilution of the 2-fold dilution that provided a Ct value in all replicates was deemed the LOD. This sample was further analysed 20 times to assess the precision of the assay with 95% confidence.

The specificity of primers and probes were theoretically assessed using NCBI nucleotide Basic Local Alignment Search Tool (BLASTn) to identify potential non-target amplification of other amoebae and protozoans that colonise the gills of marine fish. The actual specificity was assessed by testing the assays against 7 in-house amoeba cultures previously isolated from AGD-affected Atlantic salmon gills (N. perurans strain MP2, Vexillifera sp. strain MX6, Pseudoparamoeba sp. strain MX1, Vannella sp. strain MV3, MV4 and MV5 and Nolandella sp. strain MX5) (English et al. 2019) and 4 amoeba cultures obtained from American Type Culture Collection (ATCC) (N. pemaquidensis ATCC® 50172 TM, Acanthamoeba jacobsi ATCC® 30732 TM, Nolandella sp. ATCC® PRA-27 TM and Hartmannella vermiformis ATCC® 30967 TM).

The reproducibility of each assay was assessed by testing 10 gill swab DNA samples in triplicate on 3 consecutive days. These samples had previously tested positive for the target amoeba in preliminary experiments. The reproducibility was analysed by the coefficient of variation for intra- and inter-assay variation. The inter-assay differences were also assessed with a 1-way ANOVA. All qPCR validation experiments had the appropriate controls, including positive control and no-template negative control.

2.5. Inhibition

To determine whether any amplification inhibitors were present within the gill swab DNA, 5 gill swabs were diluted 2-fold 7 times such that the highest concentration was 150 ng µl⁻¹ (undiluted) and the lowest was 2.34 ng µl⁻¹ (1/64 dilution). All dilutions had a working stock volume of 10 µl. Each of the diluted samples were then spiked with 0.008 ng of plasmid DNA specific to the existing N. perurans assay (NPJ) (Downes et al. 2015). Templates (1.8 µl in 5 µl reaction) from each of the spiked dilution series were then tested in triplicate according to the qPCR protocol outlined by Downes et al. (2015).

2.6. qPCR analysis of gill swab survey

To quantify the relative abundance of several amoeba taxa on the gills of farmed Atlantic salmon, the 5 newly designed and 1 previously published qPCR assay was performed on the gill swabs collected from farm sites. Template (30 ng) DNA was tested in 5 µl reactions using MyTaq™ DNA Polymerase (Bioline) with 3 technical replicates across a 384 well PCR plate. DNA was amplified using a ViiA™ 7 Real-Time PCR Machine (Applied Biosystems) with the thermal cycling conditions specific for each assay (Table 2). A 10-fold dilution series of plasmid DNA specific to the amplicon of each assay, with a known copy number, was amplified on each plate to generate a standard curve used to determine the copy number of each sample. Each plate also included a positive control, a negative control and a negative-process control (a blank sample extracted along with the gill swab samples). An external process control (salmon elongation factor-1α) was also
run against each sample (Bruno et al. 2007). Samples were determined as positive detections if the average Ct of 2 or 3 technical replicates was lower than the limit of detection specific to each assay.

2.7. Statistical analysis

All graphics and analysis were performed in R version 3.5.2 (R Core Team 2018). To evaluate how spatial, temporal and pathological variables influenced N. perurans infection load, the N. perurans 18S rRNA copy number was first log 10 transformed after increasing all values by 1 to account for the zero values. To investigate the correlation between gill score and N. perurans abundance, a linear model was fitted. The mean N. perurans abundance with respect to time and site was compared using a 2-way ANOVA and then a Tukey post-hoc test to determine significant pairwise differences among sampling events. A paired t-test was used to compare the mean N. perurans prevalence by site and the mean gill score by site.

3. RESULTS

3.1. qPCR assays design and optimisation

After the design and optimisation of TaqMan qPCR assays specific to the chosen amoebae, the probe and primer sequences and reaction parameters were established as listed in Table 2. Results related to the primer and probe optimisation are available in Tables S1 & S2 in the Supplement at www.int-res.com/articles/suppl/q011p405_supp.pdf.

3.2. qPCR assays validation

Each newly designed qPCR assay underwent a series of validation experiments to determine their performance using both plasmid DNA and amoeba DNA as template. The results of these experiments are summarised in Table 3 with reference to the supplementary material containing the raw data. Despite NPJ being a previously published assay (Downes et al. 2015), a new standard curve and limit of detection was generated so that it was specific to our qPCR conditions (i.e. machine, taq, plasmid DNA template, etc.). Additionally, the precision, reproducibility and specificity experiments were repeated for NPJ, to act as a benchmark assay.

All 6 assays generated a linear standard curve, indicated by the coefficient of determination (R²) value close to 1. The amplification efficiency varied between the assays, with the Pse being the most efficient at 99.98% and the ParE being the least at 65.41%. The limit of detection is first presented in Table 3 as the mean Ct of the dilution which provided a Ct value for all 4 technical replicates. This value was then converted to 18S rRNA copies and shows that NPJ was the most sensitive assay, detecting up to 0.66 18S rRNA copies µl⁻¹, while ParE was the least sensitive with a limit of detection equivalent to 7871 18S rRNA copies µl⁻¹. The final dilution
Table 3. Summary of validation metrics for the 6 amoeba Taqman qPCR assays. LOD: limit of detection; CV: coefficient of variation; Nol: *Nolandella* spp.; Pse: *Pseudoparamoeba* sp.; ParE: *P. eilhardi*; VanC2: *Vannellida* species 2; VanC3: *Vannellida* species 3; NPJ: *N. perurans*

<table>
<thead>
<tr>
<th></th>
<th>Nol</th>
<th>Pse</th>
<th>ParE</th>
<th>VanC2</th>
<th>VanC3</th>
<th>NPJ</th>
<th>Supplementary reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>Standard curve R²</td>
<td>0.998</td>
<td>0.990</td>
<td>0.996</td>
<td>0.996</td>
<td>0.998</td>
<td>0.998</td>
<td>Fig. S1</td>
</tr>
<tr>
<td>Amplification efficiency (%)</td>
<td>82.23</td>
<td>99.98</td>
<td>65.41</td>
<td>72.82</td>
<td>98.28</td>
<td>99.21</td>
<td>Fig. S1</td>
</tr>
<tr>
<td>Limit of detection (Ct)</td>
<td>36.37</td>
<td>38.36</td>
<td>36.53</td>
<td>37.21</td>
<td>37.73</td>
<td>38.73</td>
<td>Table S3</td>
</tr>
<tr>
<td>Limit of detection (18S rRNA copies µl⁻¹)</td>
<td>13.50</td>
<td>1.35</td>
<td>7871</td>
<td>39.25</td>
<td>1.09</td>
<td>0.66</td>
<td>Table S3</td>
</tr>
<tr>
<td>Precision of LOD (mean ± SD)</td>
<td>36.44 ±0.57</td>
<td>39.42 ±0.58</td>
<td>35.96 ±0.57</td>
<td>37.98 ±0.62</td>
<td>35.98 ±0.62</td>
<td>39.48 ±0.54</td>
<td>Fig. S2</td>
</tr>
<tr>
<td>Intra-assay variance (mean CV %)</td>
<td>0.81</td>
<td>1.02</td>
<td>2.09</td>
<td>1.01</td>
<td>1.45</td>
<td>0.53</td>
<td>Table S4</td>
</tr>
<tr>
<td>Inter-assay variance (mean CV %)</td>
<td>1.06</td>
<td>1.35</td>
<td>2.19</td>
<td>1.31</td>
<td>1.72</td>
<td>0.88</td>
<td>Table S4</td>
</tr>
</tbody>
</table>

which determined the limit of detection was tested a further 20 times to determine the precision of the assay at a 95% confidence level. All precision data, apart from 2 outliers (Fig. S2), were close to the mean value for each assay, indicating low variability (all SD < 0.7).

The final 2 values in Table 3 refer to the reproducibility of each assay between technical replicates (intra-assay variance) and between different days (inter-assay variance) and is expressed as the coefficient of variation (CV). The mean intra-assay variation of all assays ranged between 0.53% for the NPJ assay and 2.09% for the ParE assay. In contrast, the mean inter-assay variation of all assays ranged between 0.88% for NPJ and 2.19% for ParE, which were again the best and worst performers respectively. A 1-way ANOVA further indicated there was no significant difference between the assays results conducted on 3 different days, as all p-values were >0.05 (Nol: 0.90, Pse: 0.99, ParE: 0.96, VanC2: 0.96, VanC3: 0.99, NPJ: 0.72).

Initially, all the newly designed probes and primers were deemed theoretically specific as they had no complete matches to existing reference sequences in the BLASTn database. The assays specificity was further tested against DNA samples from clonal amoeba cultures and specific plasmid DNA in the cases where no amoeba DNA-positive control was available. A positive detection was defined by a mean Ct value lower than the limit of detection and is marked in bold in Table 4. All assays appeared to be species-specific in terms of the amoeba strains against which they were tested, except for NPJ and Nol. NPJ had some cross-reactivity with *Pseudoparamoeba* sp., while Nol amplified 2 *Nolandella* species, the ATCC *Nolandella* sp. strain as well as the *Nolandella* culture it was designed to detect. The Nol assay was therefore defined as genus-specific, rather than species-specific.

To assess the presence of amplification inhibitors in gill swab DNA sourced from Tasmanian salmon farms, 5 samples were diluted, spiked with the same DNA samples and tested against each assay.

Table 4. Specificity of the 6 qPCR assays (see Table 3 for abbreviations) determined by testing each assay against various amoeba DNA samples listed as taxa (strain). Positive detections are indicated in **bold**. –: undetected
amount of plasmid DNA and then tested in triplicate. No inhibition was detected because amplification remained consistent across the dilution series (Fig. S3).

### 3.3. Survey of amoeba prevalence and abundance

Of the 6 amoeba species surveyed on the gills of farmed Atlantic salmon, *Neoparamoeba perurans* was the most prevalent (Fig. 2). *N. perurans* was dominant across both temporal and spatial factors, except at 1 site at a single time point. Atlantic salmon held at Killala during May 2017 were not colonised by detectable levels of *N. perurans*; rather, *Nolandella* spp. was highly prevalent, infecting 92% of salmon which had an average gill score of 0.225 (Fig. 2A). Except for this sampling event, *Nolandella* spp. were not detected in any other gill samples. *Pseudoparamoeba* sp. and the 2 Vannellida species were detected on the gills of a relatively small proportion of Atlantic salmon compared to *N. perurans* and *Nolandella* spp., and these few instances were not correlated with temporal or pathological factors. The remaining species, *Paramoeba eilhardi*, was not detected on any Atlantic salmon gills. In terms of the amoeba community on individual fish, non-*N. perurans* amoeba species were detected on 41 Atlantic salmon throughout the entire survey, and of these, 8 individual fish tested positive for *N. perurans* and 1 other amoeba species. There were no individual fish that tested positive for ≥3 amoeba species colonising the gills.

The relative abundance of *N. perurans* was further analysed to explore correlations between gross gill pathology, farm site and time points. The relative abundance of *N. perurans* increased with increasing gill score, indicated by the upward trend of the regression line fitted to Fig. 3A. Although this model was significant (p = 2.2 × 10⁻¹⁶), only 25% of the variability in *N. perurans* abundance could be explained by gross gill pathology (R² = 0.247). In terms of temporal and spatial factors (Fig. 3B), there were differences by site (p = 9.57 × 10⁻⁵), but when considering both, there was a significant difference between May 2017 and Aug 2017 (p = 8 × 10⁻⁶) and May 2017 and Feb 2018 (p = 0.004). The trend in *N. perurans* abundance across site and time (Fig. 3B) also reflected the fluctuations in *N. perurans* prevalence (Fig. 2), in that the number of Atlantic salmon infected with *N. perurans* increased with the *N. perurans* load on the gills. Abundance also aligned with the severity of gross gill pathology, shown by the average gill scores marked on Fig. 2.

When considering site-based differences alone, the overall *N. perurans* infection prevalence significantly differed (p = 1.374 × 10⁻⁶) with 47.4% of salmon sam-

![Fig. 2](image.png)

Fig. 2. (A–G) Prevalence of Atlantic salmon gills that were colonised by *N. perurans* (NPJ), *Nolandella* spp. (Nol), *Pseudoparamoeba* sp. (Pse) and Vannellida (VanC2, VanC3) at each farm site (Killala, Tasman) over a year. *P. eilhardi* was not included because no Atlantic salmon tested positive for the amoeba species. The average gill score (gill index [GI]) for each site and each time point is marked in the respective panels. The Feb 2018, Killala sampling event marked with ‡ is blank because no samples could be collected.
pled at Killala infected compared to 76.1% at Tasman. The abundance of \textit{N. perurans} on the gills was also significantly different between the sites (p = 2.232 × 10^{-5}), as salmon at Killala had a lower average \textit{N. perurans} load of 1.213 log10 18S rRNA copies compared to Tasman at 1.871 log10 18S rRNA copies. There was no significant difference in severity of gross gill pathology with respect to site (mean gill score Killala = 0.6, Tasman = 0.69, p = 0.366).

4. DISCUSSION

Five qPCR assays were designed to detect amoeba species previously isolated from AGD-affected Atlantic salmon \textit{Salmo salar} and then used in a spatial-temporal survey to infer the potential role these species play alongside \textit{Neoparamoeba perurans} in AGD of farmed Atlantic salmon. \textit{N. perurans} was by far the dominant species colonising Atlantic salmon gills (Fig. 2.), and their abundance was related to gross pathology (Fig. 3A), farm sites and time points throughout the year (Fig. 3B). There were very few detections of the 5 non-\textit{N. perurans} species apart from \textit{Nolandella} spp. which was dominant at 1 site (Killala) during 1 sample time (May 2017) (Fig. 2A). Significantly, this is the first molecular-based study to investigate the ecology of multiple amoeba species in association with AGD and shows that, although the gills of farmed seawater Atlantic salmon subjected to a freshwater treatment regime are colonised with a diverse Amoebozoan community (English et al. 2019), \textit{N. perurans} dominates this ecological niche, consistent with its documented role in AGD pathology (Crosbie et al. 2012).

As gross gill pathology became more severe, \textit{N. perurans} abundance was positively correlated (Fig. 3A). Although this linear relationship was expected, the amount of variation in \textit{N. perurans} abundance, expressed as 18S rRNA copies, across each gill score was not anticipated. A positive relationship between agent abundance and gross gill pathology (gill score 1 to 4) has previously been demonstrated with a small number of gill swabs from farmed Tasmanian salmon (n = 32) (Bridle et al. 2010). This previous experiment retrieved a much tighter association with an R² of –0.998, compared to the model from the present study where only 25% of the variability in \textit{N. perurans} abundance could be explained by gill pathology (R² = 0.247). The increased unexplained variability could be reflective of the use of a different \textit{N. perurans}-specific assay or variation in swabbing technique at each sampling event. It could also be because our study swabbed salmon across a number of time points at 2 different sites, as opposed to a singular sampling event, showing that \textit{N. perurans} abundance and AGD progression was responsive to some site-based and seasonal variables.

From a commercial perspective, it is useful to understand whether particular farm sites produce fish with less severe AGD. The 2 farm sites in this survey were chosen based on hypothetical differences in terms of \textit{N. perurans} infection prevalence and load due to the contrasting environmental condi-
tions and the observation that lower salinity reduces _N. perurans_ abundance (Wright et al. 2017). The Killala site is exposed to more freshwater runoff, attributed to its location in the Huon Estuary, while the Tasman farm is more oceanic (Fig. 1). Indeed, we did find Killala had an overall lower percentage of Atlantic salmon infected with _N. perurans_ and a lower mean _N. perurans_ load compared to Tasman. However, there was no difference in severity of gross gill pathology between the 2 sites, with an overall mean gill score of 0.6 at Killala and 0.69 at Tasman (p-value = 0.366). It appears that fish stocked at the Tasman lease are more likely to be infected with _N. perurans_, but these differences do not impact the severity of gross gill pathology at the 2 sites when considering gross pathology alone. However, it is likely a significant difference in AGD severity would become evident over more sampling events and with more sensitive methods for scoring pathology, such as histopathology (Adams & Nowak 2003).

Temporal differences in _N. perurans_ infection prevalence and load also differed by location. At the Tasman lease, the highest _N. perurans_ infection prevalence and load was recorded in May 2017 and Feb 2018 and the lowest in Nov 2017 (Fig. 3B). At the Killala lease, the highest was recorded in Aug 2017 and the lowest in May 2017. Although there were too few sampling events to draw conclusions on seasonal variation from our survey, the temporal trends at the Tasman lease broadly reflect past seasonal trends recorded in AGD-affected farmed Tasmania Atlantic salmon (Clark & Nowak 1999). AGD prevalence has been shown to peak in summer, followed by a second spike in autumn, and this also correlated with higher water temperature and salinity (Clark & Nowak 1999). Similarly, an _in vitro_ growth optimisation study found peak _N. perurans_ growth occurs at 15°C and a salinity of 35 ppt (Collins et al. 2019). However, the Killala lease did not reflect that higher Tasmanian water temperatures support the highest _N. perurans_ infection prevalence and load. Instead, the time point with the coolest water temperature of 11.9°C, Aug 2017, had the highest _N. perurans_ infection prevalence (92.5%) and load at this lease. This observation supports that _N. perurans_ are very adaptable to environmental parameters (Lima et al. 2016, Collins et al. 2019) and infectious over a range of temperatures and salinities (Clark & Nowak 1999).

The reasons for the dichotomy between _Nolandella_ spp. and _N. perurans_ prevalence in May 2017 (Fig. 2A,D) is speculative. One explanation could relate to exposure time to _N. perurans_. The Atlantic salmon sampled at Killala in May 2017 were transferred to sea <1 mo before sampling and had never been treated for AGD (2 to 3 h freshwater bath), while all other fish swabbed during the survey were at sea for ≥3 mo and had undergone freshwater treatments. The Killala site had also been fallowed for 7 mo prior to restocking, and it is possible that _N. perurans_ density within the environment decreased over this fallowing period. Then, in the absence of _N. perurans_, other amoeba species filled this niche. Moreover, _Nolandella_ sp. strain MX5 (English et al. 2019) showed tolerance to salinity <35 ppt during our in-house culturing. Further research could investigate whether _Nolandella_ spp. are specific to sites with more estuarine conditions or their presence is attributed to the absence of _N. perurans_.

Further research should also investigate whether the presence of _Nolandella_ spp. is either reflective of healthy Atlantic salmon gills, simply a commensal bystander or a potential player in the early stages of gill disease. For instance, a similar amoebic disease, nodular gill disease (NGD), which affects freshwater salmonids, is proposed to have a multi-amoebo aetiology, and different amoeba species are hypothesised to influence different phases of disease development (Dyková et al. 2010, Dyková & Tyml 2016). The Atlantic salmon that tested positive for _Nolandella_ spp. and negative for _N. perurans_ had an average gill score of 0.225, which was a relatively low level of AGD-like gross gill pathology compared to the other sampling events. The gross AGD-like lesions were recorded with the Taylor et al. (2009) scoring method and were not confirmed by histology because destructive sampling did not suit working alongside the commercial farm. Because histology was not used to confirm an AGD diagnosis, it is possible some gross gill lesions could be unrelated to AGD. The use of histology in conjunction with qPCR should be considered in future surveys aiming to identify relationships between specific amoeba taxa and gill pathology, or more specifically, in future investigation into the significance of _Nolandella_ spp. in relation to AGD.

_Pseudoparamoeba_ sp., the 2 Vannellida species and _Paramoeba elhardi_, are minor players in the farmed salmon gill Amoebozoa community on the east coast of Tasmania. With so few detections, no strong spatial, temporal or pathological correlations could be established for the majority of the non-Neoparamoeba species. Our findings do not corroborate an earlier study which is the only other published survey that attempted to associate AGD pathology with a variety of protozoans (Bermingham & Mulcahy 2006). The Ireland-based survey used histological observations...
to conclude that abundance of amoebae other than *Neoparamoeba* spp., *Ichthyobodo*-like flagellates and trichodinid ciliates correlated with gill pathology, while the abundance of *Neoparamoeba* spp. did not. This differs from our findings, where the abundance of *Neoparamoeba* sp. correlated with increased gross gill pathology, while the abundance of non-*Neoparamoeba* did not. Indeed, the contrasting results could be due to different locality, the possible mistaken documentation of complex gill disease rather than AGD (Herrero et al. 2018), or the study being conducted before species-specific molecular tools were available, relying instead on histology of gills. This technique is vulnerable to inaccurate documentation of amoeba community composition due to loss of amoebae during fixation, difficulty identifying protozoans in histology sections, and the limited sample size from each gill basket (1 section from 1 gill arch). Since this early study, molecular diagnostics identified *N. perurans* as the causative agent in the earliest outbreaks of AGD in Ireland which were preserved in histological samples (Downes et al. 2018).

The absence of *P. eilhardi* detections suggests either the isolation of this species documented by English et al. (2019) was a one-off occurrence or the qPCR assay was not sensitive enough. The latter reason is most likely, as the *P. eilhardi* assay performed relatively poorly in terms of amplification efficiency, limit of detections and intra- and inter-assay variance (Table 3). In comparison, the other 4 newly designed qPCR assays performed markedly better in the validation experiments and were of similar efficiency to the previously published *N. perurans* (NPJ) assay (Downes et al. 2015). The only concern with the remaining assays was that NPJ and Nol had some cross reactivity with non-target species. However, the NPJ assay only narrowly returned a positive result when tested against a high concentration of *Pseudoparamoeba* DNA, and the Nol assay was cross reactive with a species from the same genus and was thus deemed genus-specific, not species-specific. We therefore believe this lower specificity would not have affected the final conclusions. Hence, the qPCR assays, excluding perhaps the *P. eilhardi* assay, were considered fit for purpose, accurately quantifying their respective amoeba in the survey gill swabs.

5. CONCLUSION

While a diversity of amoebae colonise the gills of farmed Atlantic salmon *Salmo salar* with AGD, *Neoparamoeba perurans* is the dominant species, and increasing *N. perurans* load on the gills correlates with increasing AGD pathology. However, despite *N. perurans* being the dominant species, there were times when they could not be detected and *Nolandella* spp. were highly prevalent on gills with low levels of gross gill pathology. Further research should investigate whether *Nolandella* spp. frequently colonises farmed Atlantic salmon gills and whether they are parasitic or commensal.

Acknowledgments. The authors acknowledge Tassal Operations Pty Ltd for their support of the project, particularly Ben Morrisby, Eduardo Canosa, Christine Huynh and Bradley Evans for their technical assistance. We also thank Ben Maynard for sample collection and Megan Rigby for the valuable suggestions on the manuscript. This work was supported by CSIRO Aquaculture and The Crawford Fund (Crawford-in-Queensland 2016 Post Graduate Research Award).

LITERATURE CITED


Downes JK, Collins EM, Morrissey T, Connor IO and others


Submitted: April 2, 2019; Accepted: July 4, 2019
Proofs received from author(s): August 18, 2019