18S ribosomal DNA-based PCR identification of Neoparamoeba pemaquidensis, the agent of amoebic gill disease in sea-farmed salmonids

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ABSTRACT: Neoparamoeba pemaquidensis is a parasomal amoeboid protozoan identified as the agent of amoebic gill disease (AGD) in Atlantic salmon Salmo salar reared in sea-pens in Tasmania, Australia, and coho salmon Oncorhynchus kisutch farmed on the west coast of the USA. Outbreaks of AGD caused by immunologically cross-reactive paramoebae have also been reported in sea-farmed salmonids in several other countries. Complete 18S rDNA sequences were determined for respective paramoebae isolated from infected gills of salmon from Tasmania and Ireland, and N. pemaquidensis isolates from the USA and UK, including representative free-living isolates. Alignments over 2110 bp revealed 98.1 to 99.0% sequence similarities among isolates, confirming that paramoebae implicated in AGD in geographically distant countries were homologous and belonged to the same species, N. pemaquidensis. The results supported previous findings that N. pemaquidensis exists as a widely distributed, amphizoic marine protozoan. Partial 18S rDNA sequences were obtained for the ultrastructurally similar species, N. aestuarina, and for the morphologically similar but non-parasomal amoeba Pseudoparamoeba pagei. N. aestuarina had 95.3 to 95.7% sequence similarities with N. pemaquidensis strains, which distinguished 2 closely related but separate species. Neoparamoeba spp. were not analogous to P. pagei or to other marine Gymnamoebia. We designed 4 oligonucleotide primers based on elucidated 18S rDNA sequences and applied them to single-step and nested 2-step PCR protocols developed to identify N. pemaquidensis to the exclusion of apparently closely related and non-related protistan taxa. Nested PCR was able to detect the AGD parasite from non-purified, culture-enriched net microfouling samples from Atlantic salmon sea-pens in Tasmania, and confirmed that N. pemaquidensis was also responsible for AGD in chinook salmon O. tshawytscha in New Zealand. Our sequence and PCR analyses have now shown that AGD affecting 3 different salmonid species farmed in 4 countries are associated with N. pemaquidensis. A species-specific diagnostic PCR provides for the first time, a highly specific detection and identification assay for N. pemaquidensis that will facilitate future ecological and epidemiological studies of AGD.

KEY WORDS: Amoebic gill disease · Neoparamoeba pemaquidensis · Neoparamoeba aestuarina · 18S ribosomal DNA · PCR · Salmo salar · Salmonids

INTRODUCTION

Neoparamoeba pemaquidensis (Page, 1987) is an amoeboid protozoan that causes amoebic gill disease (AGD) in salmonids sea-farmed in several different countries (Munday et al. 2001). AGD remains the most important parasitic disease affecting sea-farmed Atlantic salmon Salmo salar L. and rainbow trout Oncorhynchus mykiss Walbaum in Tasmania, Australia (Munday et al. 1990, 2001). Outbreaks of AGD have also been reported in Atlantic salmon farmed in Chile, France (Clark & Nowak 1999), Ireland (Rodger & McArdle 1996, Palmer et al. 1997) and Spain (Munday et al. 2001), and in coho salmon O. kisutch Walbaum in
Washington State and California, USA (Kent et al. 1988), chinook salmon *O. tsawytscha* Walbaum in New Zealand (C. Anderson pers. comm.), and in rainbow and brown trout *S. trutta* L. in France (Findlay et al. 1995). The disease appears to be uniquely enzootic in southeast Tasmania, where Atlantic salmon reared in sea-pens are affected by recurring, seasonally influenced outbreaks (Clark & Nowak 1999). In contrast, AGD has only occurred as sporadic or minor outbreaks in other regions. AGD due to *Neoparamoeba* sp. have also affected non-salmonid fishes, including turbot, *Scophthalmus maximus* L. and European seabass *Dicentrarchus labrax* L. reared in seawater in Spain and the Mediterranean respectively (Dyková et al. 1995, 1998, Dyková & Novoa 2001).

*Neoparamoeba pemaquidensis* belongs to a group of exclusively marine amoebae formerly classified within the genus *Paramoeba* Schaudinn, 1896 (Cann & Page 1982, Page 1987). The paramoebae are distinguished by possession of 1 or more DNA-positive, membrane-bound inclusions located adjacent to the cell nucleus and known as parasomes. The genus *Neoparamoeba* was proposed by Page (1987) and distinguished the species *N. pemaquidensis* and *N. aestuarina* from *P. eilhardi* (type species of genus *Paramoeba*) based on cell surface structural and other morphological and ultrastructural differences. Both *N. pemaquidensis* and *N. aestuarina* are widely distributed species and have commonly been isolated as free-living trophozoites from a range of coastal marine habitats around the world (Page 1973, Cann & Page 1982). Although the 2 species have been separated based on small differences in physical and *in vitro* growth characteristics (Page 1970), there is little to differentiate them ultrastructurally (Cann & Page 1982). As a consequence, there has been some confusion as to whether *N. aestuarina* also exists in parasitic form and causes AGD similar to that associated with *N. pemaquidensis* (Dyková et al. 2000). The occurrence of *N. pemaquidensis* in both parasitic and free-living forms has indicated that this is an amphizoic species (Page 1974, Roubal et al. 1989).

AGD-affected fishes develop characteristically similar gill cytopathology, regardless of the fish species (Kent et al. 1988, Roubal et al. 1989, Dyková et al. 1998, Munday et al. 2001). Infected gill filaments exhibit epithelial hyperplasia and hypertrophy leading to complete fusion of the secondary lamellae. Infecting amoeba cells are generally abundant within the interlamellar spaces associated with the hyperplastic epithelium. Consequently, histopathological examination of gill sections has served as a reliable but non-specific method for diagnosis of AGD and for confirming the presence of infecting amoeba cells (Clark & Nowak 1999). Howard & Carson (1993) developed an indirect fluorescent antibody test (IFAT) based on polyclonal antisera raised against *Neoparamoeba pemaquidensis* isolated from AGD-affected Atlantic salmon in Tasmania. IFAT is applied to the serological screening of gill sections and mucous smears for *N. pemaquidensis*, and has been used to identify gill-associated paramoeba cells and isolates from other countries where AGD has been reported (Rodger & McArdle 1996, Palmer et al. 1997, Munday et al. 2001). IFAT remains the ‘gold standard’ test for confirming AGD cases in Tasmanian salmon farms, although an alternative high-throughput immuno-dot blot test has recently also been developed by Douglas-Helders et al. (2001). Although immunologically-based screening has proved useful in the diagnosis of AGD in infected gill samples (Zilberg & Munday 2000), its application to the specific identification of *N. pemaquidensis* has been limited by inter-species cross-reactivity of the available polyclonal antisera. In particular, antisera developed against *N. pemaquidensis* to date have cross-reacted with *N. aestuarina* and the non-parasomal amoeba *Pseudoparamoeba pagei* (Douglas-Helders et al. 2001). These species have been found in the same coastal habitats as the implicated AGD parasite (Page 1970, 1979) and have the potential to interfere with cross-specific identification tests applied to environmental samples.

Diagnostic PCR primers targeting specific regions of the 18S (small subunit) ribosomal RNA gene have frequently been used for the identification and detection of fish parasites from the host and from environmental samples (Cunningham 2002). In this study, we determined the 18S rDNA sequences of representative strains of *Neoparamoeba pemaquidensis* (including AGD-associated and free-living isolates), and of representative strains of *N. aestuarina* and *Pseudoparamoeba pagei*. Comparative 18S rDNA sequence analysis was used to clarify the relationships between AGD isolates of *N. pemaquidensis* from different countries of origin and with *N. aestuarina*, and was applied to the development of specific diagnostic oligonucleotides for *N. pemaquidensis*. The development of a species-specific nested PCR protocol useful for the identification of *N. pemaquidensis* from new AGD and free-living paramoeba isolates, and with potential application for direct detection of the parasite from fish-farm environmental samples is described.

**MATERIALS AND METHODS**

**Strains and culture maintenance.** *Neoparamoeba pemaquidensis* isolates that had been culture-purified from AGD-infected Atlantic salmon gills were used for analysis in this study (Table 1). These included 4 sepa-
rate isolates cultured from sea-penned Atlantic salmon in southeast Tasmania by T. Howard, Fish Health Unit, Department of Primary Industries Water and Environment (DPIWE), Launceston, Tasmania, and 1 isolate originating from an AGD outbreak in farmed Atlantic salmon in Ireland in 1995 (Palmer et al. 1997). Reference parasitic and free-living strains of *N. pemaquidensis*, and other species of marine amoebae including representative cultures of *N. aestuarina* and *Pseudoparamoeba pageni*, obtained from the American Type Culture Collection (ATCC), Manassas, Virginia, USA, and the Culture Collection of Algae and Protozoa (CCAP), Ambleside, Cumbria, UK, were also used for DNA analysis (Table 1). Also included were 4 amoeba isolates of uncertain taxa isolated from AGD-infected Atlantic salmon gills in Tasmania (D. Zilberg pers. comm.).

Amoeba strains were obtained as live *in vitro* cultures with the exception of the Irish *Neoparamoeba pemaquidensis* isolate AVG 8194, which was received as 70% ethanol-preserved cell suspension. Tasmanian *N. pemaquidensis* isolates were culture-cloned on malt-yeast extract seawater (MYS) agar incubated at 20°C as previously described (Page 1973, Roubal et al. 1989). Cultures were grown monoxenically with *Stenotrophomonas maltophilia* bacterium (DPIWE Strain PA1) as the food source (Douglas-Helders et al. 2001). Subcultures were propagated by excising small squares of agar containing amoeba cells and inverting them onto fresh bacteria-seeded MYS agar plates.

Table 1. *Neoparamoeba* spp. and other marine amoeba strains and isolates used for DNA analysis in this study. ATCC 50172 is an amoebic gill disease (AGD) isolate from coho salmon (Kent et al. 1988), while all other AGD isolates were from Atlantic salmon

<table>
<thead>
<tr>
<th>Species</th>
<th>Strain</th>
<th>Source</th>
<th>Origin</th>
</tr>
</thead>
<tbody>
<tr>
<td><em>Neoparamoeba pemaquidensis</em></td>
<td>ATCC 50172</td>
<td>Gills, AGD</td>
<td>Washington, USA</td>
</tr>
<tr>
<td></td>
<td>ATCC 30735</td>
<td>Free-livinga</td>
<td>Virginia, USA</td>
</tr>
<tr>
<td></td>
<td>CCAP 1560-4</td>
<td>Free-living</td>
<td>Wales, UK</td>
</tr>
<tr>
<td></td>
<td>CCAP 1560-5</td>
<td>Free-living</td>
<td>Wales, UK</td>
</tr>
<tr>
<td></td>
<td>PA 027</td>
<td>Gills, AGD</td>
<td>Tasmania, Australia</td>
</tr>
<tr>
<td></td>
<td>PA 010</td>
<td>Gills, AGD</td>
<td>Tasmania, Australia</td>
</tr>
<tr>
<td></td>
<td>PA 011</td>
<td>Gills, AGD</td>
<td>Tasmania, Australia</td>
</tr>
<tr>
<td></td>
<td>PA 111</td>
<td>Gills, AGD</td>
<td>Tasmania, Australia</td>
</tr>
<tr>
<td></td>
<td>AVG 8194</td>
<td>Gills, AGD</td>
<td>Ireland, UK</td>
</tr>
<tr>
<td><em>Neoparamoeba aestuarina</em></td>
<td>CCAP 1560-7</td>
<td>Free-living</td>
<td>Portugal</td>
</tr>
<tr>
<td><em>Pseudoparamoeba pageni</em></td>
<td>CCAP 1566-1</td>
<td>Free-living</td>
<td>England, UK</td>
</tr>
<tr>
<td>Unspeicated amoebaab</td>
<td>UP 1-6</td>
<td>Gills, AGD</td>
<td>Tasmania, Australia</td>
</tr>
<tr>
<td></td>
<td>UP 3a-4</td>
<td>Gills, AGD</td>
<td>Tasmania, Australia</td>
</tr>
<tr>
<td></td>
<td>UP 4a-2</td>
<td>Gills, AGD</td>
<td>Tasmania, Australia</td>
</tr>
<tr>
<td></td>
<td>UP 4b-5</td>
<td>Gills, AGD</td>
<td>Tasmania, Australia</td>
</tr>
</tbody>
</table>

aOriginally cultured from free-living cells in marine environments
bIdentified by 18S rDNA sequences as *Paraflabellula hoguei* (Amaral Zettler et al. 2000)
GeneQuant instructions. Purified DNA was quantified using the QIAquick PCR purification kit (Qiagen) according to the manufacturer’s instructions using spin columns from the QIAquick PCR purification kit (Qiagen Life Technologies) as molecular size marker. We used a 1 kb DNA ladder (New England Biolabs or Invitrogen Life Technologies) as molecular size marker.

Amplified 18S rDNA was purified for sequencing using spin columns from the QIAquick PCR purification kit (Qiagen) according to the manufacturer’s instructions. Purified DNA was quantified using the GeneQuantpro RNA/DNA calculator (Amersham Pharmacia Biotech). Sequencing reactions were performed according to manufacturer’s instructions using the ABI BigDye™ Terminator Cycle Sequencing Ready Reaction kit (PE Applied Biosystems) and cycle amplification performed in the GeneAmp® System 9700 thermocycler. DNA sequence data was obtained with the ABI Prism™ 377 DNA sequencer (PE Applied Biosystems). Both coding and non-coding strands of the 18S rDNA were independently sequenced and checked against the complementary strands for each amoeba strain. Overlapping sequences obtained with different primers were also compared. Sequence discrepancies were checked by repeat-sequencing of the DNA region in question using the appropriate primers.

**Sequence analysis.** Analysis and alignment of DNA sequences were performed using the Sequence Navigator™ Version 1.0.1 program (PE Applied Biosystems). Sequences generated in this study were subjected to a BLAST sequence similarity search (Altschul et al. 1990) via the National Center for Biotechnology Information (NCBI) web server (www.ncbi.nlm.nih.gov), to confirm that 18S rRNA gene sequences were obtained and to screen for potentially related taxa. Sequences from Neoparamoeba spp. and strains generated in this study were aligned and compared against each other, and also with the 18S rDNA sequences of selected Gymnamoebia (Page 1987) and other relevant protistan taxa available from the GenBank sequence database. Percentage sequence similarities between the respective Neoparamoeba spp. and strains were calculated as described by Stothard et al. (1998).

**Development of PCR primers for Neoparamoeba pemaquidensis.** Complete 18S rDNA sequences obtained for *N. pemaquidensis* were examined for suitable annealing sites for the design of species-specific oligonucleotide primers. Suitable target regions consisted of sequences that hybridise between the various *N. pemaquidensis* stains and isolates, but contain sufficient nucleotide differences to distinguish this species from other taxa. Deduced primer sequences were examined for possible complementarity with 18S rDNA of non-target amoebae (Table 1), and subjected to random BLAST sequence similarity searches to determine if homologous or related sequences were present in the existing sequence databases. Candidate primers and potential primer pairs were also checked to minimise self-complementarity. We synthesised (Genset Pacific) 4 oligonucleotides targeting suitable regions of the 18S rDNA of *N. pemaquidensis* for evaluation (Table 2); 3 forward primers (Table 2) were respectively paired with the reverse primer rNp-Hx49 and tested for amplification of *N. pemaquidensis* 18S rDNA. Optimal PCR conditions were the same for all 3 primer sets and similar to the standard conditions for 18S rDNA amplification, with the exceptions that 0.1 µM of each primer was used per reaction and the stringency of annealing was increased to 58°C during the PCR cycling steps. Direct-sequence analyses showed that only primer fNp-Hxe23b1 would be able to distinguish *N. pemaquidensis* from the closely related species, *N. aestuarina*. Primer fNp-Hxe23b1 has 7 non-complementary bases within the 24-base primer annealing site in *N. aestuarina* 18S rDNA (Table 3). In contrast, *N. aestuarina* possessed near identical sequences to *N. pemaquidensis* at the respective annealing sites of primers fNp-Hxe23a1, fNp-Hx43a1, and rNp-Hx49.

**Nested PCR protocol for Neoparamoeba pemaquidensis.** Analyses of initial non-purified (mixed) cultures obtained from environmental samples have suggested that *N. pemaquidensis* cells occur at low concentrations (Tan et al. 2002). A nested 2-step PCR procedure was developed to maximise the chances of positive detection of the AGD parasite from these samples. Total cellular DNA extracted from these samples were subjected to a primary (first-round) PCR amplification using the forward primer fNp-Hxe23a1 and the universal 18S rDNA reverse primer of Medlin et al.

<table>
<thead>
<tr>
<th>Primer</th>
<th>Sequence (5′–3′)</th>
<th>Positions</th>
</tr>
</thead>
<tbody>
<tr>
<td>fNp-Hxe23a1</td>
<td>CATCTCTTCTACTACCTTCATG</td>
<td>715–737</td>
</tr>
<tr>
<td>fNp-Hxe23b1</td>
<td>GTGATGATGATGACTCCCTACTCG</td>
<td>780–803</td>
</tr>
<tr>
<td>fNp-Hx43a1</td>
<td>GTGATGACCAAATGATTACATCCG</td>
<td>1540–1561</td>
</tr>
<tr>
<td>rNp-Hx49</td>
<td>CACAACAAACTCAGCTTACC</td>
<td>2009–2030</td>
</tr>
</tbody>
</table>
Table 3. Neoparamoeba spp. Respective 18S rDNA sequences corresponding to annealing positions of N. pemaquidensis diagnostic primer fNp-Hxe23b1. Dots indicate matching bases; non-matching bases are highlighted

<table>
<thead>
<tr>
<th>Species/primer</th>
<th>Sequence (5’–3’)</th>
</tr>
</thead>
<tbody>
<tr>
<td>N. pemaquidensis</td>
<td>GTGAGTGATGAGTAGACCTAYTG ^ ^ ^ ^ ^ ^ ^ ^</td>
</tr>
<tr>
<td>Primer fNp-Hxe23b1</td>
<td>GTGAGTGATGAGTAGACCTA T G</td>
</tr>
<tr>
<td>N. aestuarina</td>
<td>GTGAGTGATGAGTAGACCTA T G</td>
</tr>
</tbody>
</table>

*Consensus sequence for N. pemaquidensis derived from all strains sequenced*

(1988) and standard reaction and cycling conditions previously described. One µl of the ca. 1.4 kb primary PCR reaction product was used as template for a secondary (second-round) nested PCR utilising the forward primer fNp-Hxe23b1 and reverse primer rNp-Hx49 (Table 2) to amplify N. pemaquidensis 18S rDNA to the exclusion of other species. Reaction and cycling conditions (58°C annealing) for secondary amplification were similar to those already described for N. pemaquidensis-specific PCR.

Specificity of the nested PCR was tested by screening against cellular DNA extracts from the target and non-target amoebae in Table 1. Genomic DNA from Atlantic salmon was also subjected to the nested PCR to test for possible cross-reactivity, since the procedure was likely to be applied to samples associated with salmon tissues. Specificity of the secondary primer pair, rNp-Hxe23b1 and rNp-Hx49, when used in the 2-step PCR protocol, was compared to single-step PCR amplification using these primers. Efficacy of the nested PCR was evaluated using whole cells of Neoparamoeba pemaquidensis Strain PA 027 suspended in filter-sterilised, natural seawater. Duplicate DNA extractions were performed on respective serial 2-fold dilutions starting from a suspension of 2560 cells down to 40 cells, and 30 ng cellular DNA was used as the primary PCR template.

**PCR of Neoparamoeba pemaquidensis from salmon farm samples.** The N. pemaquidensis-specific nested PCR was applied to samples processed from biofouling contamination films on marine net-pens. Microfouling fractions (biofilm) on submersed pen netting were sampled from commercial Atlantic salmon farm sites in southeast Tasmania during biofouling field surveys for N. pemaquidensis (Tan et al. 2002, Douglas-Helders et al. 2003). Biofilm was swabbed from lifted near-surface netting and transferred onto MYS agar (Tan et al. 2002). Plates incubated at 20°C for 7 to 10 d were harvested, and cell suspensions were subjected to total DNA extraction and PCR analysis. The net biofilm sampled from 1 farm site formed part of a 10 wk survey to investigate the effects of antifouling paint treatments on the distribution of N. pemaquidensis on sea-pen netting (Douglas-Helders et al. 2003). Samples tested by PCR comprised culture-enriched biofilm smears collected from 5 separate marine net pens on Weeks 2 and 8 of that study. Netting from these pens had either been treated with a water-based synthetic latex (Wattyl, Australia) antifouling paint (1 pen: P1), a copper-based (Hempe Paint, Australia) antifouling paint (2 pens: P2 and P3), or left as untreated controls (2 pens: P4 and P5). In addition, 4 ethanol-preserved amoebae suspensions cultured from gills of chinook salmon Oncorhynchus tshawytscha in New Zealand were received for PCR testing. These cultures were originally derived from sea-penned fish in Marlborough Sounds, New Zealand, that were affected by an outbreak of AGD. Putative identification of a para-moeboid gill parasite had been made by gill histology and IFAT identification of the cultures (C. Anderson pers. comm.).

**Sequence verification of PCR products.** The second-round PCR products of arbitrarily selected, positive, farm-associated samples were subjected to DNA sequencing to verify that the respective amplicons were derived from Neoparamoeba pemaquidensis. Sequences were obtained using the diagnostic primers in Table 2, and universal eukaryotic 18S rDNA primers (Elwood et al. 1985) that anneal to positions internal to the expected PCR product. DNA sequences from PCR-positive samples were aligned and compared to 18S rDNA sequences of Neoparamoeba spp., and homologies were assessed.

**RESULTS**

**18S rDNA sequence analysis of marine amoebae**

PCR amplification using the universal 18S rRNA gene primers of Medlin et al. (1988) and cellular DNA from Neoparamoeba pemaquidensis strains produced 18S rDNA amplicons of ca. 2.1 kb in size. Complete 18S rDNA sequences were obtained for 4 AGD-associated N. pemaquidensis isolates from Tasmania by direct sequencing of the amplified gene products. In each case, 2109 bp of the 18S rDNA sequence were obtained for comparative analyses. All 4 Tasmanian isolates possessed identical sequences, and PA 027 was selected as the representative Tasmanian AGD strain in the present study. Respective 18S rDNA amplicons from the Irish AGD isolate, and from free-living and AGD-associated strains from the USA and UK, were similar in molecular size to that of PA 027. Complete 18S rDNA sequences of 2107 to 2110 bp
were obtained for these strains. Free-living *N. pemaquidensis* Strains CCAP 1560-4 and CCAP 1560-5 were isolated from the same region in Wales and had identical 18S rDNA sequences. All other *N. pemaquidensis* strains tested had highly similar but non-identical sequences. *N. aestuarina* Strain CCAP 1560-7 produced a ca. 2.1 kb PCR product from which a near-complete 18S rDNA sequence of 1901 bp was obtained. In comparison, amplified 18S rDNA from *Pseudoparamoeba pagei* and 4 unidentified gill amoeba isolates were smaller in size at ca. 1.8 kb. A partial gene sequence consisting of 1352 bp was obtained for *P. pagei* Strain 1566-1, while complete 18S rDNA sequences of 1861 bp were obtained for the unidentified isolates. The 4 unspeciated gill isolates shared identical 18S rDNA sequences, and comparison with available sequences in the GenBank database revealed near identical (99.9%) sequence similarity with the non-parasomal leptomyxid amoeba species *Parafflabellula hoguae* (GenBank Accession No. AF293899).

The complete 18S rDNA sequences obtained for *Neoparamoeba pemaquidensis* Strains PA 027, AVG 8194, CCAP 1560-4, CCAP 1560-5, ATCC 50172, and ATCC 30735 in this study were deposited with GenBank under Accession Nos. AF371967 through to AF371972 respectively. Partially completed 18S rDNA sequences were deposited with GenBank for *N. aestuarina* CCAP 1560-7 (Accession No. AF371973) and *Pseudoparamoeba pagei* CCAP 1566-1 (Accession No. AY277798). The complete 18S rDNA sequence obtained for putative *Parafflabellula hoguae* isolate, UP1-6 from gills of Atlantic salmon farmed in Tasmania was deposited under Accession No. AY277797.

### Sequence homogeneity among *Neoparamoeba pemaquidensis* strains

The respective 18S rRNA genes of AGD-associated *Neoparamoeba pemaquidensis* strains from Tasmania, Ireland and the USA, and reference free-living strains from the USA and UK, shared high levels of sequence homology (98.1 to 99.0% sequence similarities). These levels of sequence similarities confirmed that AGD isolates from geographically distant countries and infecting at least 2 salmonid species belonged to the same paramoeba species. Comparative 18S rDNA analysis showed that *N. aestuarina* is closely related to but a distinct species from *N. pemaquidensis*. Comparison of the 1901 bp 18S rDNA segment of *N. aestuarina* CCAP 1560-7 revealed 95.3 to 95.7% sequence similarities with analogous aligned sequences of the various *N. pemaquidensis* strains. The 18S rDNA sequences obtained for *N. pemaquidensis* and *N. aestuarina* were not homologous to any other marine Gymnamoebia or other protistan taxa when screened against the GenBank database, and represented new and unique sequence data for these species at the time of this study.

### Identification of *Neoparamoeba pemaquidensis* using diagnostic PCR primers

The candidate *Neoparamoeba pemaquidensis* diagnostic primer sets fNp-Hxe23a1/rNp-Hx49, fNp-Hxe23b1/rNp-Hx49, and fNp-Hx43a1/rNp-Hx49 each produced a specific amplification product when tested using the standard PCR protocol on cellular DNA prepared from the respective *N. pemaquidensis* strains (Fig. 1), regardless of the source or country of origin.
origin of the isolate. The expected molecular sizes of specific PCR products produced from these primer sets were ca. 1316, 1250 and 491 bp respectively. A single amplicon of the expected molecular size was produced using all primer sets, indicating that the respective primers specifically amplified the targeted 18S rDNA region from *N. pemaquidensis* DNA. In contrast, no amplification product was produced from DNA extracted from a putative *Parafflabellula hoguae* isolate cultured from Atlantic salmon gills, or from *Pseudoparamoeba pagei* (Fig. 1). In addition, none of the candidate diagnostic primers were complementary to other potentially occurring marine organisms when subjected to BLAST analyses.

**Specificity and efficacy of Neoparamoeba pemaquidensis nested PCR**

Nested *Neoparamoeba pemaquidensis*-specific PCR utilising 2 separate primer sets was found to be specific for all strains of *N. pemaquidensis*, while not cross-reacting with DNA from non-target organisms (Fig. 2).

The nested PCR did not cross-react with the closely related species *N. aestuarina*, nor with the non-parasomal but morphologically similar *Pseudoparamoeba pagei*. Specificity-testing of the nested PCR showed no amplification of *N. aestuarina* with the *N. pemaquidensis*-specific primer set fNp-Hxe23b1/rNp-Hx49, while cross-specific amplification occurred using primer set fNp-Hx43a1/rNp-Hx49 (Fig. 2). This confirmed that primers fNp-Hx43a1 and rNp-Hx49 annealed with *N. aestuarina* 18S rDNA, while primer fNp-Hxe23b1 did not, as predicted by direct-sequence analyses. None of the diagnostic primers reacted with Atlantic salmon DNA, allowing PCR detection of the AGD parasite in samples associated with fish tissues without interference from host DNA. No variation in specificity was observed between single-step PCR amplification and secondary amplification in the nested 2-step PCR procedure using the *N. pemaquidensis*-specific primer set fNp-Hxe23b1/rNp-Hx49 (Fig. 2). The nested PCR was able to detect down to ca. 40 *N. pemaquidensis* cells suspended in sterile seawater with confidence (Fig. 3), but lower cell numbers were not tested.
Replicate biofilm cultures from each of 5 net-pens (Pens P1 to P5) were obtained at Weeks 2 and 8 of a 10 wk pen-netting antifouling-treatment study. PCR was used to screen 25 cultured biofilm samples consisting of 5 samples from each pen at Week 2 and 15 cultured biofilm samples consisting of 3 samples from each pen at Week 8 (Fig. 4). *Neoparamoeba pemaquidensis* was detected by PCR in net biofilm sampled from all antifouling treated net-pens (P1 to P3) and from 1 untreated control pen, P5, at both Weeks 2 and 8 of the study. In contrast, *N. pemaquidensis* was not detected by PCR in any of the samples from the second untreated control pen, P4. The total percentage PCR-positive samples obtained for net-pens treated with water-based synthetic latex antifoulant (1 pen: P1), copper-based antifoulant (2 pens: P2 and P3), and untreated pens (2 pens: P4 and P5) were 40, 50 and 20% respectively at Week 2; and 33, 33 and 17% respectively at Week 8. In addition to the antifouling treatment study, 15 other net biofilm cultures from miscellaneous sea-pens located at a different Tasmanian farm site were subjected to DNA analysis; of these 53% were PCR-positive for *N. pemaquidensis*. Furthermore, 2 of the ethanol-preserved amoeba culture suspensions prepared from chinook salmon gills in New Zealand were positive for *N. pemaquidensis* following amplification by nested PCR.

**Sequence confirmation of PCR-positive samples**

DNA sequences were obtained for nested PCR products amplified from 2 randomly selected culture-enriched biofilm samples from Atlantic salmon sea-pens in Tasmania and 2 chinook salmon gill cultures from New Zealand. The DNA sequences of amplicons obtained from net microfouling biofilms were homologous with the 18S rDNA of *Neoparamoeba pemaquidensis*, with 99.0% sequence similarity to Tasmanian AGD Strain PA 027 when 1212 bases were aligned and compared. Sequences from both biofilm samples

![Fig. 3. *Neoparamoeba pemaquidensis*. Detection of Strain PA 027 cells suspended in filter-sterilised seawater by species-specific, nested 2-step PCR. Respective cellular DNA templates were extracted from dilutions containing 2560 cells (Lane 1), 1280 cells (Lane 2), 640 cells (Lane 3), 320 cells (Lane 4), 160 cells (Lane 5), 80 cells (Lane 6), and 40 cells (Lane 7). Control reactions included amplification of 30 ng cellular DNA prepared from cells harvested directly from agar culture of Strain PA 027 (Lane 8), and a no-template control (Lane 9). Arrow indicates position of 1250 bp PCR product. Lanes M: Invitrogen 1 kb DNA ladder](image)

![Fig. 4. *Neoparamoeba pemaquidensis*-specific nested PCR of enriched biofilm (microfouling) samples collected from netting of 5 sea-pens at Atlantic salmon farm site in Tasmania, during 10 wk antifouling paint treatment trial. Netting from sampled pens had either been treated with water-based synthetic latex antifouling paint (Pen P1), a copper-based antifouling paint (Pens P2 and P3), or were untreated controls (Pens P4 and P5). Biofilm samples from same net-pens (P1 to P5) collected at Week 2 (5 samples per net-pen) and Week 8 (3 samples per net-pen) of the trial were tested by PCR. Control reactions included amplification of 30 ng cellular DNA prepared from cells harvested directly from agar cultures of *N. pemaquidensis* Strains ATCC 50152 and PA 027 (Lanes +), and a no-template control (Lane –). Large arrow indicates position of expected ca. 1250 bp PCR product; small white arrows indicate PCR-positive samples. Lane M: Invitrogen 1 kb DNA ladder](image)
were identical. Similarly, amplicon sequences from chinook salmon gill cultures were homologous with *N. pemaquidensis* PA 0127 18S rDNA, with 98.4% sequence similarity over 925 bases. However, the true extent of sequence divergence between *N. pemaquidensis* strains in the farm sample cultures and index AGD strains in Table 1 could not be determined since sequences from the entire 18S rDNA were not available for comparison.

**DISCUSSION**

The parasitic agent of AGD in sea-farmed Atlantic salmon in Tasmania was found by morphological and ultrastructural comparisons to most closely resemble *Neoparamoeba pemaquidensis* (Kent et al. 1988, Roubal et al. 1989, Munday et al. 1990). Outbreaks of histopathologically identical AGD have also affected sea-farmed salmonids in the USA and several European countries (Kent et al. 1988, Palmer et al. 1997, Munday et al. 2001). Although paramoebae from these cases were immunologically reactive with antisera prepared from Tasmanian isolates, there remained insufficient evidence to conclude that the various AGD isolates belonged to the same species. Using 18S rDNA sequence data, we have now confirmed that AGD isolates of paramoeba from Tasmania are homologous and identical to the species *N. pemaquidensis*. The Tasmanian isolates possessed greater than 98% sequence similarity with both parasitic and free-living isolates of *N. pemaquidensis* originating from the USA and UK. Sequence homogeneity among strains isolated from different countries in 3 continents also supported previous findings that *N. pemaquidensis* is a widely distributed marine species (Page 1973).

The genus *Neoparamoeba* currently includes 2 species, *N. pemaquidensis* and *N. aestuarina* (Page 1970, 1987). Free-living trophozoites of both species have been found in similar marine environments and are morphologically and ultrastructurally similar, making cells difficult to differentiate (Page 1970, Cann & Page 1982). In the current study, analysis of 18S rDNA sequences was able to distinguish *N. pemaquidensis* and *N. aestuarina* as 2 separate but closely related species. The levels of sequence similarities among a diverse collection of *N. pemaquidensis* strains (maximum sequence dissimilarity of 1.9%) were consistent with a single homologous sequence type suggested for putative definition of species within another gymnamoeboid genus, *Acanthamoeba* (Stothard et al. 1998). In contrast, *N. aestuarina* had at least 4.3% sequence dissimilarity with strains of *N. pemaquidensis*. The *Neoparamoeba* spp. were not homologous to any other protistan taxa that presently exist in the eukaryotic 18S rDNA sequence database. Recent preliminary phylogenetic analyses based on 18S rDNA sequence data suggest that *N. pemaquidensis* and *N. aestuarina* represent a distinct clade independent of other marine Gymnamoebia (Fiala & Dyková 2003), including the morphologically similar but non-parasomal amoeba *Pseudoparamoeba pagei* (Page 1987, Elliott et al. 2001). A specific detection method for *Neoparamoeba pemaquidensis* is required for ecological and environmental studies, since the AGD-causing organism is closely related to *N. aestuarina* and possibly other as yet uncharacterised members of the genus. Currently available diagnostic methods for the AGD parasite are mainly based on immunodetection, either by IFAT or immuno-dot blot (Howard & Carson 1993, Douglas-Helders et al. 2001). However, polyclonal antisera developed for *N. pemaquidensis* cross-react with *N. aestuarina* and *Pseudoparamoeba pagei* (Douglas-Helders et al. 2001), although not with a range of other more distantly related amoebae found co-associated with gills of AGD-affected fishes (Howard & Carson 1993). This cross-specificity may not be important for detection of *N. pemaquidensis* in heavily infected gill samples or from culture cloned isolates, but presents a potential limitation to detection in non-purified cultures or environmental samples where higher numbers of immunologically related organisms may be present. Analysis of 18S rDNA sequences has enabled us to develop a diagnostic PCR protocol specific for *N. pemaquidensis*.

The 18S rRNA genes of some eukaryotic taxa contain unique expansion sequence segments that are highly variable between phylogenetically divergent or non-related organisms (Stothard et al. 1998, Sims et al. 1999). These sequence segments are situated in non-conserved regions of the gene and are often responsible for the variations in nucleotide length observed between 18S rDNA of divergent organisms. The design of 4 oligonucleotide PCR primers for *Neoparamoeba pemaquidensis* was based on sequence sites located within 18S rDNA expansion segments observed in 2 current species of the genus *Neoparamoeba*. These non-conserved gene segments were shared by closely related *N. pemaquidensis* and *N. aestuarina*, but were absent from or very different in all other representative taxa examined. Consequently, 3 of the primers had respective annealing sites present in both *Neoparamoeba* species. However, we were able to identify 1 primer site in *N. pemaquidensis* 18S rDNA that allowed differentiation from *N. aestuarina*. PCR utilising a forward primer corresponding to this annealing site together with a co-specific reverse primer was shown to specifically amplify *N. pemaquidensis* DNA to the exclusion of *N. aestuarina*. None
of the diagnostic primers cross-reacted with other taxa used for PCR-testing.

While single-step PCR using the species-specific primer set could successfully identify Neoparamoeba pemaquidensis cells from culture-purified isolates, we also developed a nested 2-step PCR protocol for potential application to non-purified cultures from field samples, or directly from non-enriched environmental samples. The protocol relied on a semi-specific primary amplification step followed by secondary amplification using a nested *N. pemaquidensis*-specific primer set. The semi-specific, first-round primers were selected to maximise the concentration of target DNA template for the secondary PCR step while minimising possible masking of target DNA by excessive accumulation of non-target 18S rDNA expected from samples containing mixed protistan and other eukaryotic fauna. Second-round PCR essentially confirmed that *N. pemaquidensis* 18S rDNA was amplified in the first round, as well as increasing the sensitivity of detection for the target species by cumulative PCR cycling (Barlough et al. 1995). *N. pemaquidensis*-specific PCR did not amplify DNA of the non-parasomal marine gymnomoebae *Pseudoparamoeba pagei* (Page 1979) or Atlantic salmon gill isolates identified by 18S rDNA sequences in this study as *Paralabellula hoguae* (Amaral Zettler et al. 2000). The ability of the diagnostic PCR to detect *N. pemaquidensis* to the exclusion of non-related amoebae is important, since the latter may form part of a complex amoeboid fauna previously detected in gills of AGD-infected salmon reared in Tasmanian sea-pens (Howard & Carson 1992).

The threshold of PCR detection was not determined for a particular sample type in the present study, although a strong amplification product was consistently obtained for ca. 40 Neoparamoeba pemaquidensis cells seeded into sterilised seawater and subjected to DNA extraction. Suspensions harvested from enrichment cultures would be expected to contain considerably larger numbers of amoeba cells (Douglas-Helders et al. 2001). Nested PCR was able to detect the presence of *N. pemaquidensis* in culture-enriched net biofilm (microfouling) samples collected from 2 Atlantic salmon farm sites in southeast Tasmania. This was consistent with positive IFAT detection of *N. pemaquidensis* in cultured net biofouling samples by Tan et al. (2002), suggesting that pen-netting might be a significant environmental reservoir for AGD. Net-pen surfaces and associated biofouling may provide favourable substrates for the attachment of *N. pemaquidensis* cells, a prerequisite for population growth (Martin 1985). Biofilm samples collected from net-pens that had been respectively treated with 2 types of commercial antifouling paint had an overall higher incidence of *N. pemaquidensis* than samples from untreated pens. The effects of antifoulants on the dynamics and composition of net-fouling organisms appear to contribute positively to the distribution of *N. pemaquidensis* in that environment (Douglas-Helders et al. 2003). PCR detection of *N. pemaquidensis* prevalence in respective cultured biofilm samples from untreated nets and those treated with a copper-based antifoulant correlated well with results obtained by IFAT screening of equivalent but independently processed samples (Douglas-Helders et al. 2003). Good agreement between these unrelated tests served to validate the positive detection of *N. pemaquidensis* by both PCR and IFAT, and suggested that immunologically cross-reactive amoebae may be absent or occurred in negligible numbers in the net biofouling cultures. Direct validation of the nested PCR products by DNA-sequencing further confirmed that *N. pemaquidensis* was specifically detected in the net biofilm samples.

The ability to distinguish between *Neoparamoeba pemaquidensis* and *N. aestuarina* is an important consideration for future ecological and pathological studies of AGD. Morphological and ultrastructural comparisons were unable to distinguish *N. aestuarina* Strain CCAP 1560-7, from paramoebae that caused AGD in non-salmonid fish, turbot and European seabass (Dyková et al. 1998, 2000). In contrast, nested PCR was able to confirm that paramoeba responsible for AGD in New Zealand chinook salmon was *N. pemaquidensis*, demonstrating its usefulness for the specific identification of the AGD parasite. We have thus shown conclusively in this study that AGD affecting 3 different salmonid species sea-farmed in 4 countries, namely the USA (coho salmon), New Zealand (chinook salmon), Ireland and Australia (Atlantic salmon), were associated with the ubiquitous paramoeba species *N. pemaquidensis*. Whether all cases of AGD are due to *N. pemaquidensis*, or can also be caused by other parasitic *Neoparamoeba* species, is unclear (Dyková et al. 2000, Fiala & Dyková 2003). Recent analysis of new 18S rDNA sequence data has indicated that a related *Neoparamoeba* sp. distinct from both *N. pemaquidensis* and *N. aestuarina* may be associated with some cases of AGD in seawater-reared turbot in Spain (Fiala & Dyková 2003). Paramoeba isolates from related and future AGD cases could be subjected to identity confirmation using the *N. pemaquidensis*-specific PCR described in this study. To date, *N. aestuarina* has only been reported as free-living trophozoites and has not been associated with disease in fishes.

Although DNA-based detection of *Neoparamoeba pemaquidensis* has mainly been applied to culture-enriched field samples in this study, the PCR assay could also be potentially optimised for direct detection in various environmental samples without prior enrich-
ment. We have detected *N. pemaquidensis* directly in macerated net biofouling organisms, including a common bryozoan species, using our nested PCR protocol (data not shown). Tan et al. (2002) also detected the presence of paramoeba on a number of macrofouling species growing on Tasmanian Atlantic salmon pens using IFAT on culture-enriched samples. Important factors that influence the efficacy of PCR detection in crude samples include distribution of the target organism and the effects of possible inhibitors to DNA extraction and PCR amplification within the sample matrix (Hiney 1997). As such, field surveys for *N. pemaquidensis* by PCR should involve a systematic approach to sampling, since choice of appropriate DNA extraction strategies and validation of PCR results are dependent on each sample type and the inherent matrix examined (Hiney 1997, Cunningham 2002). The continuing severe impacts of AGD on the sea-farming of Atlantic salmon in Tasmania have led to an increasing interest in assessing the distribution and ecology of *N. pemaquidensis* in the associated marine environment, with the aim of developing better farm-management strategies for controlling the disease. Recent studies have investigated the distribution of paramoeba cells remaining in the gills of dead salmon (Douglas-Helders et al. 2000), in wild-caught indigenous fish species (Douglas-Helders et al. 2002), and in biofilm and macrofouling organisms associated with pen-netting as well as in the surrounding water column (Tan et al. 2002). The diagnostic PCR developed in the present study provides, for the first time, a highly specific detection and identification method for *N. pemaquidensis* that will facilitate future related ecological and epidemiological studies of AGD.

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