Monitoring *Piscirickettsia salmonis* by denaturant gel electrophoresis and competitive PCR

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ABSTRACT: Reported strains of *Piscirickettsia salmonis*, a pathogen of salmonid fishes, were analyzed by amplifying part of the internal transcribed spacer (ITS) of the ribosomal RNA (rRNA) operon followed by denaturing gradient gel electrophoresis (DGGE) of the amplicons. All amplified fragments differing in sequence were distinguished by migration during DGGE. A simpler format, constant denaturant gel electrophoresis (CDGE), allowed the same diagnostic distinctions among strains. Sampling during 1997 and 1998 of salmonids from 5 different sites on and near Chiloé Island in southern Chile displaying piscirickettsiosis revealed only *P. salmonis* resembling LF-89, the type strain first isolated in 1989. These observations are encouraging for control strategies, which might otherwise be compromised by unpredictable shifts of *P. salmonis* types in salmon farms. A competitive PCR assay offered insight about the power of PCR for quantification and about specific tissue invasiveness by this intracellular pathogen. This approach revealed that the PCR could amplify approximately 1 to 10 *P. salmonis* genome equivalents against a background of >99.9% salmonid DNA. It also raised the possibility that the salmonid brain is an important site for *P. salmonis* survival, with its bacterial load in 1 individual having been about 100 times the loads observed in liver and kidney. Pathogen detection by competitive PCR in a surface seawater sample from a netpen in use indicated a density of about 3000 to 4000 *P. salmonis* cells (or their DNA remnants) l⁻¹. Such quantitative estimates should aid decisions about disease prevention and management as, for example, choice of netpen sites following fallow periods and certification of ova, which are known conduits of infection.

KEY WORDS: *Piscirickettsia salmonis* · Genetic diversity · DGGE · CDGE · Competitive PCR

INTRODUCTION

In the late 1980s salmon aquaculture in the fiords of southern Chile faced the emergence of an infectious disease of unknown origin. Symptoms of the disease initially reported in coho salmon *Oncorhynchus kisutch* were evident after fish underwent transfer to salt water (Bravo & Campos 1989). A microbial agent was isolated (Fryer et al. 1990, Cvitanich et al. 1991) and further characterized as a novel intracellular organism, *Piscirickettsia salmonis* (type strain LF-89 = ATCC VR 1361), belonging to the gamma subdivision of the proteobacteria (Fryer et al. 1992, Fryer & Mauel 1997, Mauel et al. 1999). The agent survived in salt water and less well in fresh water (Lannan & Fryer 1994); it was horizontally transmissible by contact and cohabitation in both environments (Cvitanich et al. 1991, Almendras et al. 1997), with 1 observation of vertical transmission having been reported (Almendras & Fuentealba 1997). Susception of intermediate vectors in the transmission of *P. salmonis* has been raised (Almendras & Fuentealba 1997). Stress resulting from husbandry practices otherwise considered normal and from environmental conditions has been highlighted as important in the manifestation of the disease (Almendras & Fuentealba 1997, Olsen et al. 1997).

Piscirickettsiosis or salmonid rickettsial septicemia soon became evident in other salmonids (Atlantic salmon *Salmo salar* and rainbow trout *Oncorhynchus mykiss*) in both the northern and southern hemispheres (Fryer & Lannan 1996, Almendras & Fuentealba 1997, Olsen et al. 1997, Jones et al. 1998). The
microbial etiology of these outbreaks has been confirmed by determination of close phylogenetic affiliations to *Piscirickettsia salmonis* LF-89 (Mauel et al. 1996, 1999), serological criteria (Olsen et al. 1997) and comparative biochemistry of the microbial antigens (Jones et al. 1998). However, severe epizootics of *P. salmonis* occur only in the fish farms of southern Chile, while outbreaks of the organism in other parts of the world have been less pronounced and sporadic (for reviews, see Fryer & Lannan 1996, House et al. 1999). Other rickettsia-like organisms, different from *P. salmonis* but causing similar disease symptoms, have become evident in Chilean aquaculture (Cvitanich et al. 1995, V. Henriquez, C.O. & S.M. unpubl.), but there are as yet no molecular genetic data to shed light on their origin.

We have exploited amplification of the internal transcribed spacer (ITS) region of the ribosomal RNA (rRNA) operon of *Piscirickettsia salmonis* (Marshall et al. 1998) with 3 objectives in mind: (1) to ascertain further the degree of variation existing in this genetic region among the *P. salmonis* strains or lineages reported in the literature; (2) to establish a molecular assay that could readily monitor the lineages of the pathogen that emerge seasonally in the salmon farms in Chile; and (3) to develop an assay for assessing the number of genome copies of the microbe in environmental samples as well as in different tissues of the host during infection episodes. The first 2 objectives were accomplished by electrophoresis in denaturing polyacrylamide gels (Fischer & Lerman 1983, Hovig et al. 1991), in which PCR amplicons differing in sequence migrate distinct distances in the presence of denaturants as a result of differences in DNA stability. The third objective was accomplished by competitive PCR (Gilliland et al. 1990), which involves co-amplification of the target of interest and known quantities of a mimic fragment amplifiable by the same primer pair but yielding a product of a different length.

**MATERIALS AND METHODS**

**Bacterial strains and natural samples.** DNA extracts from *Piscirickettsia salmonis* LF-89 and EM-90 (the latter isolated in Chile from Atlantic salmon) came from the lab collection (Marshall et al. 1998). DNA samples from *P. salmonis* strains SLGO-94 (rainbow trout) and C1-95 (coho salmon), both from Chile, as well as from ATL-4-91 (British Columbia, Canada, netpen-raised Atlantic salmon) and NOR-92 (Norway, netpen-raised Atlantic salmon) were donated by M. J. Mauel. These strains are listed by Mauel et al. (1996, 1999) and House et al. (1999) along with additional information on their biological and institutional origins.

Visibly infected tissues of coho salmon and rainbow trout affected with piscirickettiosis were received from 5 different salmon farms and 1 unknown location on and near Chiloe Island (encompassing the sea to the east of the island) in the 1997 and 1998 seasons and kept frozen in RNase buffer (Marshall et al. 1998). Sampling of surface seawater (4 to 5 l) from netpens in the Caucabue Channel in the Ancud Gulf, northeast of Chiloe Island, was done essentially as by Massana et al. (1997). Prefiltration of the water was through a Whatman GF/A glass microfiber filter (1.6 µm pore size, 47 mm diameter, Clifton, New Jersey, USA), and the picoplankton remaining in the filtrate was collected on a Sterivex filtration device (0.22 µm pore size, Millipore, Bedford, Massachusetts, USA) by means of a peristaltic pump. The GF/A filter was changed 2 to 3 times per sample to permit timely filtration of the desired volumes through the Sterivex device. Hose lines were washed thoroughly with tap water between filtrations. Immediately after picoplankton collection, storage buffer (1.8 ml, 40 mM EDTA, 50 mM Tris-HCl [pH 8.0], 0.75 M sucrose) was added to the Sterivex chamber (which includes the nonremovable Millipore filter) and the chamber was then kept at -20°C until nucleic acid extraction.

**DNA extraction and PCR amplification.** DNA was released from tissues by the Chelex method and 1 µl used in PCR (total reaction volume, 12.5 µl) with primer pairs RTS1/RTS2 or RTS1/RTS4 complementary to the ITS as described previously (Marshall et al. 1998). For samples to be evaluated by competitive PCR, biopsies of <0.25 mg were collected at the end of 200 µl pipette tips and submerged in 100 µl of the Chelex bead suspension.

Extraction of nucleic acids from samples collected with Sterivex devices was initiated by the addition of lysozyme (1 mg ml-1 final concentration) to the 1.8 ml of solution in the storage chamber. This step was followed by Proteinase K (0.5 mg ml-1) digestion in the presence of 0.9% sodium dodecyl sulfate, phenol/chloroform/isoamyl alcohol extraction and ethanol precipitation. Several extraction blanks were included. Final DNA fractions were diluted to 5 ng µl-1 and 1 µl was used in PCR with primer pair RTS1/RTS4. Extraction blanks were diluted to the same degree of dilution applied to the DNA fraction from the natural samples, in order to control for any inhibitory substances in these blanks.

GC-clamped amplicons for denaturing gradient and constant denaturant gel electrophoresis (DGGE/CDE) analyses were obtained from RTS1/RTS4 amplicons by reamplification (starting with 2 µl of a 200-fold dilution of the PCR products in a total reaction volume of 25 µl) with the same primer pair except that RTS1 had a 40 bp GC ‘clamp’ added at the 5'-end (Myers et al.
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1989, Murray et al. 1996). GC-RTS1 (65 bases) had the following sequence, with the ‘clamp’ being to the left of the slash: 5'-CGCCCGCCGCCGCCGCCGCCGCCGTCCGGGCCGCCGCCGCC/GAGATAGTA-3'.

**DNA sequencing.** Double-stranded PCR products (generated with primers RTS1 and RTS4) were purified with carboxylated magnetic beads (DeAngelis et al. 1995) or on silica matrix filters (Qiagen, Valencia, CA, USA) prior to the cycle sequencing reaction. Cycling parameters applied to 30–40 ng of amplicon for 50 cycles were: denaturation, 96°C, 10 s; annealing, 66°C (found to be optimal for sequence quality), 5 s; extension, 60°C, 4 min. The ABI Prism Ready Reaction dye-terminator cycle sequencing kit was used with half the recommended volume (AmpliTaq DNA polymerase FS, PE Applied Biosystems, Foster City, CA, USA). These amplification conditions (volume, molar quantity of template and cycle number) led to 99.7% accuracy of base identification with plasmid pGEM3Z(f+) and the forward –21 M13 primer, the sequencing control included in the dye-terminator kit. Similar accuracies were seen under these conditions with a PCR product that had been exhaustively sequenced (R. Zebell & C.O. unpubl.).

An ABI PRISM model 377 DNA sequencer (PE Applied Biosystems) was used with DNA Sequence Analysis software version 2.1.2 for electrophoresis and sequence display. Sequence alignment and editing were performed with Sequencher 3.0 software (Gene Codes Corporation, Ann Arbor, Michigan, USA). Manual editing of ambiguities in the sequences produced by the instrument was performed using the rules concerning peak height patterns obtained with AmpliTaq DNA polymerase FS dye-terminator sequencing (Parker et al. 1996). The first 60 to 70 bp displayed by the pherograms (next to each sequencing primer) were not included in the analyses as they are subject to severe noise from dye-terminator ‘clouds’ (ABI Prism DNA Sequencing, Chemistry Guide, PE Applied Biosystems).

**DGGE and CDGE.** The ‘melting temperatures’ of representative PCR products were first obtained on a perpendicicular denaturing gradient gel cast with a Bio-Rad model 475 gradient former in 16 cm long plates, part of the DCode™ mutation detection device (Bio-Rad, Hercules, CA, USA). The gel of 6% 37.5:1 acrylamide/bis-acrylamide gradient (100% = 40.0 ml deionized formamide and 42.0 g urea added to 100 ml) was prepared from components in the DCode electrophoresis reagent kit for DGGE/CDGE as described by the manufacturer (Bio-Rad). One hundred μl of the PCR product was layered evenly along the top of the gel, and electrophoresis was carried out at 165 V and 56°C for 3.5 h. Subsequent parallel DGGE runs were performed with a gradient of 22 to 25% denaturants. The CDGE format, defined at a 23.5% level of denaturants, became the method of choice for monitoring natural specimens.

Gels were stained with ethidium bromide (2 μg ml⁻¹ in 1x TAE buffer) for 10 min, followed by destaining for 10 min in 1x TAE. Migration patterns were revealed by UV-fluorescence, digitized with the Gel Doc™ 1000 system (Bio-Rad), and analyzed as well as optimized for contrast and brightness with the Multi-Analyst software provided with the system. Images were saved in JPEG format and then manipulated for contrast and brightness, labeled and printed in Adobe Photoshop® 4.0. Theoretical melting curves based on sequences of amplicons were generated with the WinMelt software, PC-Windows (Bio-Rad).

**Mimic PCR.** The composite primers and the competitor fragments for the quantitative PCR were constructed using the PCR Mimic Construction kit by Clontech® Laboratories (Palo Alto, California, USA). The sequence (provided by Clontech®) of the 574 bp BamHI/EcoRI fragment of the v-erbB avian blastoma virus, from which competitor fragments are constructed, was consulted to choose positions for the ends of the composite primers. Individual composite primers were designed with the sequences of the *Piscirickettsia salmonis* primers RTS1, RTS2 and RTS4 at their 5’ ends (underlined below) and the appropriate sequences of the v-erbB template at their 3' ends. Primers RTS1mimic1 (5'-TGATTTTATTGTTTAGTGAAAAATGACGCAAGTGAAATCTCCTCCG-3'), RTS2mimic3 (5'-AAATAACCCTAAATTAATCAAGGATTTCATCTCCTCGTATAACA-3'), and RTS4mimic6 (5'-ATGCACTTATTCACTTGATCATAATTTGATTCGACCATGGC-3') were paired in the PCR as RTS1mimic1/RTS2mimic3 and RTS1mimic1/RTS4mimic6 to give amplicons from the v-erbB template of expected sizes 248 bp (157 bp longer than the RTS1/RTS2 *P. salmonis* amplicon) and 554 bp (270 bp longer than the RTS1/RTS4 *P. salmonis* amplicon), respectively.

Amplification conditions in a total reaction volume of 50 μl were as follows: 1x PCR buffer [10 mM Tris-HCl [pH 8.3], 50 mM KCl, 1.5 mM MgCl₂, and 0.001% [wt/vol] gelatin (PE Applied Biosystems)], each of the 4 deoxynucleoside triphosphates at a concentration of 0.2 mM, each oligonucleotide primer at a concentration of 0.4 μM, 0.5 ng of template, and 0.04 U of AmpliTaq DNA polymerase (PE Applied Biosystems) μl⁻¹. A hot start (D’Aquila et al. 1991) was used to impede formation of primer duplexes: a mixture containing both primers and the 4 deoxynucleoside triphosphates was added after the other components for the reaction had reached 85°C. The following thermal cycling regime was then applied: 7 cycles of denaturation at 94°C for
Fig. 1 Variation in the ITS region of the rRNA operon in reported *Piscirickettsia salmonis* strains. The polymorphic sites are listed vertically across the top, with their numbers corresponding to the LF-89 U36943.1 sequence in GenBank; lowercase letters denote sites added 3' of the indicated nucleotide relative to this sequence. The sequence we determined for LF-89 is shown at all sites; the bases in the other strains, except ATL-4-91, are unsequenced near primer RTS4. In the numbering, strains except ATL-4-91 (unsequenced near primer RTS4) an extra T in the reported sequence of a section of the ITS locus of *P. salmonis* (U36943.1, we sequenced, with no ambiguities, positions 224–458 (the entire segment between primers RTS1 and RTS4) for LF-89 and EM-90, 294–451 for SLGO-94, 252–457 for C1-95, 242–447 for ATL-4-91 and 245–455 for NOR-92 by incubation at 65°C for 1 h. The eluates (1 μl) served as templates for amplification (in a reaction volume of 50 μl) with the corresponding *Piscirickettsia salmonis*-specific primer pairs, i.e., RTS1/RTS4 and RTS1/RTS2, to reach concentrations suitable for subsequent competitive PCR experiments. Each primer was present at a concentration of 0.25 μM, the polymerase at 0.03 U μl⁻¹, and the buffer and dNTPs at the concentrations given above. The cycling regime was as detailed above except that 16 (not 18) cycles with an annealing temperature of 55°C (not 60°C) were included. Amplicons were purified on MagBeads (DeAngelis et al. 1995) and stored and diluted in 1x LTE containing 10 μg of ultrapure glycogen ml⁻¹ to prevent adhesion of nucleic acids to plastic surfaces. After electrophoresis through 1.5% agarose, quantification of the Mimic fragments was done by densitometry of stained gels using the Bio-Rad Gel Doc™ 1000 system set to show saturated pixels so as to ensure that detection remained in the linear range for adequate quantification. Special attention was given to accuracy of micropipetting and prevention of cross-contamination for competitive PCR measurements. Co-amplification of *P. salmonis* plus Mimic templates was carried out in 12.5 μl volumes with the PCR protocol described by Marshall et al. (1998).

Quantitative estimation of *Piscirickettsia salmonis* with the competitive PCR method was used also to evaluate the representation of bacterial sequences in a DNA extract from *P. salmonis* EM-90 propagated in chinook salmon Oncorhynchus tshawytscha embryo cell line CHSE-214 (Lannan et al. 1984, Marshall et al. 1998). In an attempt to reduce appreciably contributions of host DNA, the extraction was performed from a cell culture exhibiting full cytopathic effect that had been treated with DNaseI followed by washes and differential centrifugation before lysis of the bacterial cells, as outlined previously (Marshall et al. 1998).

RESULTS

Sequencing of the ITS from reported *Piscirickettsia salmonis* strains

An earlier discrepancy (Marshall et al. 1998) with the reported sequence of a section of the ITS locus of *Piscirickettsia salmonis* LF-89 prompted us to examine the ITS sequence of the other reported strains of *P. salmonis*, as noted in a preliminary account (Heath et al. 1999). We found in all *P. salmonis* strains except ATL-4-91 (unsequenced near primer RTS4) an extra T in the run of 4 Ts reported in GenBank accession U36943.1 as present in LF-89 at positions 448 to 451 (Fig. 1). A recent publication (Mauel et al. 1999) and updated

<table>
<thead>
<tr>
<th>Strain</th>
<th>Sequence 1</th>
<th>Sequence 2</th>
</tr>
</thead>
<tbody>
<tr>
<td>LF-89</td>
<td>TCA...T.AGG..TTA?</td>
<td>TCA...T.AGG..TTA?</td>
</tr>
<tr>
<td>LF-89</td>
<td>GB1 C.G. ATAK</td>
<td>C.G. ATAK</td>
</tr>
<tr>
<td>LF-89</td>
<td>M K</td>
<td>K</td>
</tr>
<tr>
<td>LF-89</td>
<td>GB2 K</td>
<td>K</td>
</tr>
<tr>
<td>SLGO-94</td>
<td>?............</td>
<td>?............</td>
</tr>
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<td>SLGO-94</td>
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<td>C......TAK</td>
</tr>
<tr>
<td>SLGO-94</td>
<td>M C......ATAK</td>
<td>C......ATAK</td>
</tr>
<tr>
<td>SLGO-94</td>
<td>GB2 C......ATAK</td>
<td>C......ATAK</td>
</tr>
<tr>
<td>C1-95</td>
<td>.......... K</td>
<td>.......... K</td>
</tr>
<tr>
<td>C1-95</td>
<td>GB1 C......AK</td>
<td>C......AK</td>
</tr>
<tr>
<td>C1-95</td>
<td>M K</td>
<td>K</td>
</tr>
<tr>
<td>C1-95</td>
<td>GB2 C......AK</td>
<td>C......AK</td>
</tr>
<tr>
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<td>.......AT.GG.....ATA?</td>
<td>.......AT.GG.....ATA?</td>
</tr>
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<td>......GAT.GG......ATA-</td>
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<td>......GAT.GG......ATAK</td>
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</tr>
<tr>
<td>NOR-92</td>
<td>M ......GAT.GG......ATAK</td>
<td>......GAT.GG......ATAK</td>
</tr>
<tr>
<td>NOR-92</td>
<td>GB2 ......GAT.GG......ATAK</td>
<td>......GAT.GG......ATAK</td>
</tr>
</tbody>
</table>
Table 1. Pairwise sequence differences in the ITS region among *Piscirickettsia salmonis* strains. Above the diagonal are the observed nucleotide substitutions, exclusive of insertions and deletions, among the sequences determined in the present study (Fig. 1). Below the diagonal are the percent sequence differences for 202 nucleotides compared among all 4 strains listed here (Fig. 1). LF-89 represents also the identical sequences found for SLGO-94 and Cl-95.

<table>
<thead>
<tr>
<th>Strain</th>
<th>LF-89</th>
<th>ATL-4-91</th>
<th>NOR-92</th>
<th>EM-90</th>
</tr>
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<tbody>
<tr>
<td>LF-89</td>
<td></td>
<td>5</td>
<td>6</td>
<td>9</td>
</tr>
<tr>
<td>ATL-4-91</td>
<td>2.5</td>
<td></td>
<td>1</td>
<td>11</td>
</tr>
<tr>
<td>NOR-92</td>
<td>3.0</td>
<td>0.5</td>
<td></td>
<td>12</td>
</tr>
<tr>
<td>EM-90</td>
<td>4.5</td>
<td>5.4</td>
<td>5.9</td>
<td></td>
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</tbody>
</table>

sequences submitted to GenBank in 2000 showed a K at 450a for all 6 strains plus 3 or 4 other differences within the segment bounded by RTS1 and RTS4 relative to the sequences Mauel and colleagues submitted to GenBank in 1995 and 1996 (Fig. 1). We noted additional differences from all the hitherto reported sequences (Fig. 1), such that fragments flanked by primer binding sites RTS1 and RTS4 were reduced from 6 to 4 sequence types (Fig. 1, Table 1), including 1 type common to Chilean isolates LF-89, SLGO-94 and C1-95 and also to *P. salmonis* sequenced from 2 Chilean rainbow trout by Marshall et al. (1998). GenBank accession numbers are AF212834 to AF212839 for the sequences we determined from the 6 reported *P. salmonis* strains and AF212840 to AF212841 for the *P. salmonis* sequences from the 2 additional Chilean fish. For a length of 236 to 237 bp, we detected 16 polymorphic sites among our own sequences (Fig. 1). Pairwise nucleotide substitutions among our sequences range from 0.5 to 5.9% (Table 1).

**Denaturant gel electrophoresis assessment of *Piscirickettsia salmonis* variation**

An initial examination of the melting properties of the RTS1/RTS4 fragment was conducted with amplicons generated from one of the most distantly related pairs, LF-89 and EM-90 (Table 1). A perpendicular DGGE analysis at 56°C revealed 2 melting transitions, at inflection points of 24 and 27% denaturant concentration (Heath et al. 1999). The theoretical melting curves for the 2 sequences had 2 melting domains, one at 62°C and a second between 62 and 63.5°C (Heath et al. 1999). Therefore, the experiment and melting theory (Lerman & Silverstein 1987) roughly agree, as the differential in the observed inflection points corresponds to 0.3°C/1% denaturant (Abrams & Stanton 1992). The prediction of higher melting values for EM-90 compared to LF-89 was confirmed in the parallel

DGGE format with a gradient ranging between 20 and 25% (results not shown) and also by CDGE (Fig. 2).

Adequate separation of the *Piscirickettsia salmonis* sequence types depended on a narrow denaturant gradient and therefore was susceptible to the vagaries of gradient casting, particularly at the beginning of the gradient, which would influence relative migration patterns and make comparisons between runs more challenging. A constant denaturant concentration of 23.5% and a 4% polyacrylamide gel gave the best separation among the sequence variants (Fig. 2). CDGE became the preferred format given the likelihood of higher reproducibility because gels are cast at a single urea-formamide concentration. Four migration classes clearly emerged: EM-90 > NOR-92 > ATL-4-91 > LF-89 = C1-95 = SLGO-94 (Fig. 2). Clustering of the last 3 agrees with our sequence data (Fig. 1). Artificial mixtures of sequence types ATL-4-91 and LF-89 were used as markers and also as a test of the technical quality of the run in subsequent assessments via CDGE (Fig. 3, lanes 1, 8 and 14). Fragments that differed by 1 substitution (Table 1) were clearly distinguished in the case of ATL-4-91 and NOR-92 (Fig. 2, lanes 4 and 6). As is evident from Fig. 2 and Table 1, the degree of difference in migration in DGGE and CDGE is not proportional to the degree of sequence difference because the least stable part of the molecule is largely responsible for retardation during migration under denaturing conditions, with other parts of the sequence and the overall length of the fragment having little influence on retardation (Fischer & Lerman 1983). Thus the nature and location of sequence differences, rather than their number, will determine the relative mobilities of variants. It should be pointed out, furthermore, that all polymorphic sites reported (from positions 293 to 450a; see Fig. 1) fall into a section that extends from
base 135 of the GC-clamped PCR fragment to its end, well within the first melting domain predicted by theory (Heath et al. 1999).

Screening of *Piscirickettsia salmonis* extracted from coho salmon (38 individuals) and rainbow trout (6 individuals) from farms in the south of Chile during the 1997 and 1998 seasons by CDGE (Fig. 3, Table 2) implied that 1 sequence type, LF-89-like, was common to all 44 samples from tissue biopsies obtained from 5 (or 6) different salmon farms on and near Chiloe Island.

### Quantitative PCR: validation and application for *Piscirickettsia salmonis*

Assessment of the efficiency of amplification of the competitor fragments synthesized for quantitative PCR by serial dilution experiments showed that both the RTS1/RTS2 and the RTS1/RTS4 Mimic fragments could be amplified from as little as approximately a single copy of template (lane 8 of Fig. 4; Heath 1999).

Reamplification of the genuine RTS1/RTS2 and RTS1/RTS4 fragments from *Piscirickettsia salmonis* LF-89 was equally efficient (Heath 1999). In addition, co-amplification of equal numbers of copies through a representative range of 10-fold dilutions of RTS1/RTS2 and its Mimic fragment also manifested similar efficiencies for both fragments in all instances (Heath 1999). The accuracy of quantification using controlled target numbers was assessed by submitting 100 copies of the *P. salmonis* RTS1/RTS2 fragment to quantitative PCR with a dilution series of its Mimic fragment. After allowing for a 2.7-fold factor in fluorescent signal intensity (248 bp Mimic fragment/91 bp *P. salmonis* fragment), the competitive measurement predicted close to 100 copies for the fragment in question (lane 5 in Fig. 4).

The effect of DNA strand length on quantification was also assessed. Evaluation of copy number from genomic templates raises questions of possibly unequal influences on amplification efficiencies (of genomic vs Mimic sequences) due to the dynamics of sep-

### Table 2. Salmonids surveyed for *Piscirickettsia salmonis* by PCR and CDGE

<table>
<thead>
<tr>
<th>Group</th>
<th>Location</th>
<th>Date</th>
<th>Salmonid</th>
<th>Number</th>
<th>Water</th>
<th>Tissue(s)</th>
<th>Type (no.)</th>
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<td>1</td>
<td>Unknown</td>
<td>Jul 1997</td>
<td>Adult cs</td>
<td>8</td>
<td>Salt</td>
<td>Kidney</td>
<td>LF-89 (8)</td>
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<td>2</td>
<td>Caucahué</td>
<td>Jul 1997</td>
<td>Adult cs</td>
<td>9</td>
<td>Salt</td>
<td>Kidney</td>
<td>LF-89 (9)</td>
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<tr>
<td>3</td>
<td>Compu</td>
<td>Jul 1997</td>
<td>Adult rt</td>
<td>1</td>
<td>Salt</td>
<td>Kidney</td>
<td>LF-89 (1)</td>
</tr>
<tr>
<td>4</td>
<td>Puerto Montt</td>
<td>Oct 1997</td>
<td>Adult cs</td>
<td>9</td>
<td>Salt</td>
<td>Kidney</td>
<td>LF-89 (9)</td>
</tr>
<tr>
<td>5</td>
<td>Quemchí</td>
<td>Oct 1997</td>
<td>Adult rt</td>
<td>3</td>
<td>Salt</td>
<td>2K, 1L</td>
<td>LF-89 (3)</td>
</tr>
<tr>
<td>6</td>
<td>Quemchí</td>
<td>Oct 1997</td>
<td>Adult cs</td>
<td>4</td>
<td>Salt</td>
<td>1K, 3L</td>
<td>LF-89 (4)</td>
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<td>7</td>
<td>Potemún</td>
<td>Oct 1997</td>
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<td>Fresh</td>
<td>Liver</td>
<td>LF-89 (2)</td>
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<tr>
<td>8</td>
<td>Potemún</td>
<td>Oct 1997</td>
<td>Smolt rt</td>
<td>2</td>
<td>Fresh</td>
<td>1B, 1L</td>
<td>LF-89 (2)</td>
</tr>
<tr>
<td>9</td>
<td>Caucahué</td>
<td>Nov 1998</td>
<td>Adult cs</td>
<td>6</td>
<td>Salt</td>
<td>Kidney</td>
<td>LF-89 (6)</td>
</tr>
</tbody>
</table>
Fig. 4. Quantification by competitive PCR of 100 copies of Piscirickettsia salmonis RTS1/RTS2 amplicon (91 bp) with RTS1/RTS2 Mimic fragment (248 bp). Lanes 2–8 show the products obtained in competitive PCR starting from 100 copies of the P. salmonis RTS1/RTS2 amplicon plus going from lane 2 to lane 8—10,000, 1,000, 500, 100, 50, 10 and 1 copies of the RTS1/RTS2 Mimic fragment. A negative PCR control is in lane 9. Lane 1 contained a 100 bp ladder with the 800 bp fragment intensified; sizes (in base pairs) are indicated on the left.

Fig. 5. Quantitative PCR with primer pair RTS1/RTS4 and DNA from (A) brain and (B) liver biopsies of an adult coho salmon with piscirickettsiosis. Chelex-extracted salmon DNA (1 μl, lanes 1–7) was amplified along with 10⁶, 10⁵, 10⁴, 10³, 10², 10¹ and 10⁰ copies (lanes 1–7, respectively) of RTS1/RTS4 Mimic fragment (554 bp). Lane 8 (B) was a negative control with no DNA added. L contains the 100 bp ladder described in Fig. 4.
DISCUSSION

Assessment of genetic variation of *Piscirickettsia salmonis* via the ITS

The ITS section of the rRNA operon is often considered the most promising locus for exploring interspecific distinctions among prokaryotes (for a recent review, see Mauel et al. 1999), though this approach is complicated when the organism harbors more than 1 rRNA operon. *Escherichia coli* has 7 such operons, with 3 containing 2 tRNA genes within their ITS region and the other 4 operons containing 1 tRNA gene different from the former 2 (Condon et al. 1995). Examples of 3 different versions of the ITS within a cell, with different numbers of tRNA insertions (3, 2 and 1) in the spacer, have been reported in the genus *Vibrio* (Chun et al. 1999). Mauel et al. (1999) have postulated, based on their cloning studies, the existence of only 1 ITS copy per genome for each of the reported *Piscirickettsia salmonis* isolates studied here. A detailed analysis of percent sequence identity is now available for 3 genetic regions of the rRNA operon in *P. salmonis* (Mauel et al. 1999). With only 1 exception (ATL-4-91 vs NOR-92), the ITS (309 nucleotide positions not including ambiguous positions) displayed at least as much variation as the 16S and 23S rRNA genes (Mauel et al. 1999). Therefore, focusing on the ITS appears justified for future surveys of genetic variation of *P. salmonis* that might alert us to the emergence of new genetic lineages of the organism in the habitats of southern Chile. The sequence discrepancies summarized in Fig. 1 do not detract from the argument that ITS is a suitable locus for such appraisals. More global assessments of the *P. salmonis* genome for genetic variation — for example by genomic fingerprinting using arbitrarily primed PCR (Welsh & McClelland 1993) — are not yet possible given the considerable obstacles in purifying the microorganism from cellular debris, including nucleic acids of its host (Barnes et al. 1998, S.M. unpubl.). Such purification is a prerequisite also for the establishment of future genome libraries representative of this intracellular pathogen.

Integration of the current studies (1997 and 1998 seasons) with an earlier one (1996 and 1997 seasons: Marshall et al. 1998) points to homogeneity of genetic types in *Piscirickettsia salmonis* obtained from coho salmon and rainbow trout from southern Chile, whether at their freshwater or salt water life stages (Fig. 3, Table 2). Earlier isolates SLGO-94 (rainbow trout) and C1-95 (coho salmon) also share an identical ITS sequence with LF-89, according to our studies (Fig. 1). Therefore, in the decade since the piscirickettsiosis syndrome was described (Bravo & Campos 1989) and *P. salmonis* strain LF-89 isolated (Fryer et al. 1990, Cvitanich et al. 1991) and typed (Fryer et al. 1992), this particular lineage has dominated the epidemiology of the disease throughout the huge expansion of the salmon farming industry in Chile. Intense habitat adaptation by *P. salmonis* might be involved. The 3 geographic isolates — LF-89 (Chile), ATL-4-91 (British Columbia, Canada) and NOR-92 (Norway) — which also comprise distinct phylogenetic isolates (Heath 1999, Mauel et al. 1999), display different degrees of virulence in controlled infections, with LF-89 being the most virulent (House et al. 1999). This picture is compatible with Chile being the site of highest incidence and prevalence of piscirickettsiosis.

Future epidemiological surveys by, for example, the DGGE protocol presented here (Fig. 3) might alter this view of the apparent constancy of genetic lineage of the *Piscirickettsia salmonis* pathogen in the Chilean salmon industry. (Preliminary reports [S. Valdebenito & B. Jauriguiberry pers. comm. from 1995 mentioned by Olsen et al. 1997, Casanova et al. 1999] have suggested that 1 to 3 additional variants of *P. salmonis*, beyond LF-89-like and EM-90, may occur among Chilean salmonids.) To date we have obtained DNA sequences for only 2 recent Chilean specimens, and they were identical to our LF-89 sequence (Marshall et al. 1998). Surveys reveal that mutation detection via DGGE is highly comprehensive (>95%; see Cotton 1997) when performed with GC-clamped fragments, provided that primers are positioned such that mutations internal to the primers occur in the first melting domain as predicted by first principles (Lerman & Silverstein 1987) and computational simulations. Therefore, new variants in the ITS region of *P. salmonis* are likely to be evident in the future via CDGE, if their sequence changes continue to occur in the section corresponding to the lowest ranking melting domain of the amplified fragment.

A number of studies have illustrated the power of denaturant gel electrophoresis to distinguish even transversions leading to the interchange of bases across strands, as would be the case for G-C and A-T transversions (Fischer & Lerman 1983, Ridanpää et al. 1995, Hepburn & Miller 1998). Judicious PCR primer positioning can lead to detection of all transversions of the type discussed here without resorting to the more elaborate analysis of heteroduplexed entities (Hepburn & Miller 1998). Therefore, there are grounds for confidence that the paucity of types as revealed by CDGE is the manifestation of sequences identical to LF-89 for the ITS region we studied. The suggestion that the same LF-89-like lineage of *Piscirickettsia salmonis* seems to have prevailed since the dawn of the salmon industry in Chile in 1982 to the present has reassuring implications in regard to the dependability of future vaccines against *P. salmonis* for coho salmon and rainbow trout.
The quantitative approach for the understanding of *Piscirickettsia salmonis*

Competitive PCR with the *Piscirickettsia salmonis* system offers a robust approach for the quantitative measurement of genome copy number of the microorganism. Results appeared to be reproducible despite different storage times and features of template quality that are otherwise hard to control, such as template lengths of genomic preparations. Confidence of measurement was well within an order of magnitude (for example, Fig. 4). Competitive PCR also allowed us to estimate the sensitivity of the current PCR assay: detection of approximately 1 to 10 *P. salmonis* genome equivalents is possible against a large background (>99%) of the salmonid DNA (for example, Fig. 5). Though simple in execution and reagents, compared to more automated and costly approaches to quantitative PCR via real-time or kinetic amplification protocols (Higuchi & Watson 1999), competitive PCR as described here is demanding in regard to accuracy of pipetting and avoidance of cross-contamination.

Obtaining precise numbers of *Piscirickettsia salmonis* genomes at different anatomical sites of infection should shed new light on the routes of invasion of the pathogen. The almost indiscriminate systemic invasion of *P. salmonis* has been well documented. In addition to characteristic lesions in the liver, kidney and spleen, the intestine, brain, skeletal muscle, heart and skin as well as macrophages have all been implicated (see Almendras & Fuentealba 1997). Nevertheless, it was surprising to discover that in one case the estimated number of resident *P. salmonis* cells in biopsies of <0.25 mg was 100 times higher in the brain compared to liver (Fig. 5). These observations raise the possibility of lower bacterial loads in tissues considered to be primary foci of invasion, compared to the brain, and suggest that this organ might be a preferred site of *P. salmonis* infection. They also point to the need for large-scale quantitative surveys of bacterial burdens in the brain and a variety of other tissues within individual fish. Non-PCR technology for quantifying the number of *P. salmonis* cells accurately in a routine fashion, even using antibodies (as in the fluorescent antibody test), is notoriously inadequate. These complications reflect the fact that this organism is an intracellular pathogen that cannot yet be grown in pure culture. We have endeavored to standardize tissue weights to a reasonable extent by taking biopsies of the same size. Furthermore, the nature of the Mimic PCR approach controls for differential efficiencies of amplification to be expected from different tissues or extracts.

A conservative calculation (1 ng for the weight of a eukaryotic cell) leads to bacterial loads of about 1 *Piscirickettsia salmonis* per 5 brain cells of the fish in the case we studied. Of course, our measurements could mean, for example, that only 1 of every 50 fish brain cells is infected, with each infected cell carrying a burden of 10 bacterial organisms. Phase contrast microscopy on *P. salmonis*-infected CHSE-214 cells suggests that at 2 to 3 d post inoculation such loads per cell are possible; counting of *P. salmonis* particles within membrane-bound cytoplasmic vacuoles reveals at least 18 organisms per vacuole in one case (Fryer et al. 1992) and in the other a minimum of 8 per vacuole (Garces et al. 1991). Similar burdens are clearly visible by transmission electron microscopy of an infected hepatocyte in *Salmo salar* (Olsen et al. 1997).

Competitive PCR should also be helpful in the elucidation of ecological scenarios in the transmission of *Piscirickettsia salmonis* leading to natural outbreaks. Its detection and quantification in seawater from a netpen at a density of 3440 cell equivalents 1⁻¹ add to earlier evidence for possible horizontal transmission via seawater (Cvitanich et al. 1991, Lannan & Fryer 1994, Almendras et al. 1997). Other avenues of transmission (see Almendras & Fuentealba 1997, Olsen et al. 1997), such as by coprophagy, via algal blooms and zooplankton (particularly copepods) and by an external hematophagous isopod parasite of salmonids (Garces et al. 1994), could be better evaluated with quantitative appraisals of bacterial loads. Certification of ova and of sites for salmon cultivation following fallow periods (Cassigoli 1994) could also benefit from the quantitative approach.

It is conceivable that amplification of DNA or even rRNA might result in false pictures, as positive signals from such targets can be obtained from nonviable bacteria (see Sheridan et al. 1998, Cook & Lynch 1999) and bacterial cell debris (Dupray et al. 1997). Reverse transcription (RT) PCR to detect bacterial messenger (m) RNAs of short half lives is increasingly being employed to monitor for the presence of viable bacteria (Sheridan et al. 1998, Cook & Lynch 1999). Competitive PCR can be implemented as part of RT-PCR (Gilliland et al. 1990). However, awareness of a viable but nonculturable state that bacteria enter upon exposure to low nutrient conditions (Steinert et al. 1997), without loss of 16S rRNA and presumably with drops in mRNA, should temper decisions to exclude DNA-based PCR assays in favor of the RT-PCR mode in evaluations of microorganisms in natural habitats.

**Molecular approaches in the management of disease in aquaculture**

Simplifications to the methodology presented here are conceivable, so as to permit more practical contributions of the molecular approach to disease manage-
ment in aquaculture of salmonids. The CDGE format is amenable to preparation of pre-cast gels in large quantities (for subsequent long-term storage). The use of fluorescent-labeled primers would eliminate staining with ethidium bromide after the electrophoresis run. For the Mimic procedure, microtiter plates with wells previously loaded with dilution series of the Mimic fragment and dehydrated for storage can be used. Pre-casting gels and delivering nucleic acids in a dehydrated state for their quantitative recovery in biochemical reactions are already practiced in today’s biotechnology commerce. Their application to aquaculture management appears as a desirable and practical goal.

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