Real-time quantitative polymerase chain reaction (QPCR) to identify Myxobolus cerebralis in rainbow trout Oncorhynchus mykiss

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ABSTRACT: This study describes the development of a TaqMan real-time quantitative polymerase chain reaction (QPCR) technique using the heat-shock protein 70 (Hsp 70) and 18S ribosomal DNA (18S rDNA) sequences to identify Myxobolus cerebralis and attempt to quantify infection severity within rainbow trout fry Oncorhynchus mykiss. Rainbow trout for this study were exposed to M. cerebralis under natural river conditions and examined for infection by histology, polymerase chain reaction (PCR) and QPCR analysis at 900 Celsius temperature units (CTUs) following exposure. Detection sensitivity by QPCR was shown to be equal to traditional PCR but greater than histopathology. Primer/probe combinations developed for this study were capable of specifically detecting M. cerebralis DNA in infected fish tissue and single triactinomyxon (TAM) spores with a sensitivity of 12.5 and 6.3 pg µl−1 of DNA for the Hsp 70 and 18S rDNA sequences, respectively. A strong relationship between the QPCR and infection severity was found for the Hsp 70 probe when parasite copy number and histology scores of 0–4 were compared (R² = 0.96, p = 0.003). However, a reduction in copy number was observed at higher histology scores for the 18S probe (scores of 4 and 5) and the Hsp 70 probe (score of 5). The results of this study demonstrate that QPCR analysis is an effective tool for detecting M. cerebralis in fish tissue and may provide a relative indication of infection severity.

KEY WORDS: Myxobolus cerebralis · Quantitative polymerase chain reaction · Histopathology · Hsp 70 · 18S rDNA

INTRODUCTION

Widespread distribution (Hoffman 1990, Bartholomew & Reno 2002) and reduced survival in wild rainbow trout Oncorhynchus mykiss associated with whirling disease (Walker & Nehring 1995, Vincent 1996) have increased the need to rapidly diagnose infections by Myxobolus cerebralis. The principal methods used to identify the parasite include pepsin-trypsin digest (PTD) (Markiw & Wolf 1974), histopathology (Lorz & Amandi 1994, Hedrick et al. 1999, Baldwin et al. 2000 as modified by Andree et al. 2002) and polymerase chain reaction (PCR) analysis (Andree et al. 1998, Schisler et al. 2001, Baldwin & Myklebust 2002). Histopathology and PTD analysis can also be used to determine infection severity, but both techniques lack sensitivity for detecting low-level infections. Although myxospore enumeration, as determined from PTD analysis, represents a relative level of infection, spore proliferation and virulence of M. cerebralis infection can be influenced by environmental conditions and host life-stages; therefore this technique is typically not used as the primary measure of severity (Nehring & Walker 1996). In addition, peri-
ods of 3 to 11 mo may be required to allow spore development to occur and PTD analysis to be performed (Andree et al. 2002).

PCR analysis is currently the most sensitive method available for detecting Myxobolus cerebralis, but infection intensity cannot be estimated by PCR. Although effective for clinical diagnosis, the above procedures can be time-consuming, labor-intensive and subjective in their ability to distinguish M. cerebralis from other closely related myxozoan species. Based on this and a need to further develop rapid diagnostic capabilities, we initiated a study to determine if a real-time quantitative polymerase chain reaction (QPCR) assay would be an effective method for detecting M. cerebralis DNA within infected fishes and quantifying infection severity during early myxospore development.

QPCR assays have been established for a number of fish pathogens (Overturf et al. 2001, del Cerro et al. 2002, Bilodeau et al. 2003, Corbeil et al. 2003) and offer a number of potential advantages. This technique is similar to traditional PCR in that it requires a series of short oligonucleotide primers for amplification of target sequences. However, in addition to dual primer sets, QPCR utilizes a sequence-specific fluorescent probe. During polymerization, this probe is cleaved, resulting in a level of fluorescence that can be quantified as a relative measure of template material when compared to a known standard. Fluorescence can then be visualized and measured using computer-based programs, therefore eliminating the need for the post-gel visualization required for traditional PCR techniques.

Molecular markers selected for PCR-based assays generally target gene sequences that are highly conserved and species-specific. The current PCR-based assay for Myxobolus cerebralis targets a ribosomal sequence (Andree et al. 1998, Kelley et al. 2004), but a specific assay based on the heat-shock protein 70 (Hsp 70) gene of this parasite has recently been developed (Epp et al. 2002). Targeting ribosomal genes is common since they are found in all living systems and control the synthesis of protein from messenger RNA (mRNA) (Hillis & Dixon 1991). The highly conserved nature of these genes is useful for species identification (Hillis & Dixon 1991), and PCR will typically yield a high degree of sensitivity associated with the presence of several sequence copies/genome (Bialek et al. 2000). Another gene that is highly conserved and important in living organisms is the heat-shock protein gene. Heat-shock proteins function as molecular chaperones that bind to other proteins and assist with folding and native confirmation required for protein-protein interactions (Feder & Hofmann 1999). These genes are often used as phylogenetic indicators to distinguish between large groups of organisms (Feder & Hofmann 1999), and multiple copies per genome are generally present. Therefore, the Hsp 70 and 18S ribosomal DNA (18S rDNA) genes are ideal candidates for development of a QPCR analysis for M. cerebralis.

As a fish becomes infected and the etiological agent begins to replicate within its host, a pathogen’s genetic copy or DNA concentration should increase as infection progresses. By using gene-specific probes, QPCR analysis should be capable of detecting increasing pathogen loads or possible carrier states within the host. This has been suggested for a number of fish pathogens including infectious haematopoietic necrosis virus (IHNV), Flavobacterium psychrophilum and Edwardsiella ictaluri (Overturf et al. 2001, del Cerro et al. 2002, Bilodeau et al. 2003). Potential benefits from a QPCR assay for Myxobolus cerebralis include sensitivity associated with PCR methodology, reduced labor, and potentially cost. A combined analysis may be capable of determining the presence and severity of infection within a single analysis. Reduced labor and cost could also be associated with the potential to examine infected fish tissues prior to myxospore development, ultimately eliminating the need for extensive rearing periods following parasite exposure in laboratory-based studies. Therefore, the objectives of this study were to (1) design and test primer/probe combinations specific for M. cerebralis based on the Hsp 70 and 18S rDNA gene sequences, (2) determine specificity and sensitivity of the QPCR technique in relation to current diagnostic methods, and (3) identify potential relationships, during early stages of infection, between the parasite’s genetic copy number and infection intensity within rainbow trout.

MATERIALS AND METHODS

QPCR primer/probe design and instrumentation. A partial 581 base-pair nucleotide sequence from the Hsp 70 gene (Accession No. AY553906) for Myxobolus cerebralis was isolated and characterized by Pisces Molecular LCC using PCR amplification with degenerate primers (Epp et al. 2002). Dr. J. Wood provided this Hsp 70 nucleotide sequence, and the 18S rDNA gene sequence was obtained from GenBank (Accession No. U96492). Primer/probe combinations from both sequences were designed using Primer Express Software version 1.5 (Applied Biosystems). A BLAST search (Altschul et al. 1997) to identify homology of primer/probe combinations with other genetic sequences within the GenBank database was performed to determine potential for non-specific amplification. Primers were synthesized by Integrated DNA Technologies and probes were constructed by Applied Biosystems.
**Hsp 70.** Forward 414F, 5'-CGAAGAACTCAATG-GCTGATCTT-3' and reverse 471R, 5'-TTGGTCAA-TTTGCTACAGTAA-3' primers were designed to amplify an 80 base-pair region of the Hsp 70 sequence for *Myxobolus cerebralis*. The TaqMan probe 448P, 5'-TCTTGACCCCGTTGAGAA-3' was designed with the fluorescent reporter 6-carboxyfluorescein (FAM-6) linked to the 5’ end and a proprietary (Applied Biosystems) non-fluorescent quencher (NFQ) attached to a minor groove-binding molecule (MGB) which was linked to the 3’ end.

Prior to analysis, all samples were standardized to 20 ng µl⁻¹ total DNA using a Perkin Elmer MBA 2000 spectrophotometer. Reaction conditions for the Hsp 70 sequence were established to provide optimal amplification and were thereafter performed as follows: reaction volume was 20 µl and contained 1× TaqMan universal master mix (Applied Biosystems), 0.6 µM forward primer, 0.6 µM reverse primer, 0.8 µM fluorescent labeled probe, and 20 ng µl⁻¹ DNA. The 20 µl reaction was transferred to individual wells of a 96-well optical plate (Applied Biosystems), capped, and placed in a BioRad iCycler real-time PCR detection system. Duplicate wells of no template controls (NTC) were incorporated into each 96-well plate to test for cross contamination between samples. The amplification profile consisted of initial denaturing at 95°C for 10 min, followed by 45 cycles of 15 s at 95°C and 1 min at 60°C. All samples were again analyzed in duplicate to demonstrate repeatability.

**18S rDNA.** Forward 811F, 5'-TGAATAAATCAGAGTGCTAAAGC-3' and reverse 937R, 5'-TTGGTGCTGTATGCTGTAACTG-3' primers were designed to amplify a 140 base-pair region of the 18S rDNA sequence for *Myxobolus cerebralis*. The TaqMan probe 888P, 5'-TGTGACAAATAGCG-3' was designed with the same fluorescent reporter and quencher system described above. Sequence positions within the Hsp 70 and 18S rDNA genes are shown in Table 1.

Reactions for the 18S rDNA sequence were established to provide optimal conditions for amplification and were thereafter performed as follows: reaction volume was 20 µl and contained 1× TaqMan universal master mix (Applied Biosystems), 0.15 µM forward primer, 0.15 µM reverse primer, 0.2 µM fluorescent labeled probe, and 20 ng µl⁻¹ DNA. The 20 µl samples were prepared and inserted into the iCycler detection system as previously described. The amplification profile consisted of initial denaturing at 95°C for 10 min, followed by 40 cycles of 15 s at 95°C and 1 min at 62°C. All samples were again analyzed in duplicate to demonstrate repeatability.

**PCR amplification with QPCR primer combinations.** Primer combinations for the Hsp 70 and 18S rDNA sequences were first examined by conventional PCR analysis. Reaction volumes were 25 µl and contained 1× Qiagen PCR buffer, 2.5 mM MgCl₂, 0.6 µm dNTPs, 2U Taq polymerase, and 0.6 µM of each primer. Gradient conditions followed previously established protocols (Baldwin & Myklebust 2002) and amplification was performed using a PTC-100 thermal cycler (MJ Research).

**Standard design.** The amplified region of the Hsp 70 gene was cloned into the pGEM®-T Easy Vector (Promega) and the amplified region of the 18S rDNA gene was cloned into the pCR®II-TOPO® Vector using the Zero Blunt®TOPO®PCR Cloning Kit (Invitrogen) following the manufacturers’ protocols. Inserts were sequenced at Washington State University’s DNA sequencing laboratory (Pullman) using an Applied Biosystems 3100 genetic analyzer to confirm presence of appropriate inserts. The nucleotide length of each plasmid was counted and concentration was determined by OD₂₆₀ using a Perkin Elmer MBA 2000 spectrophotometer. The values obtained were used to determine the relationship between the concentration of plasmid DNA and its copy number, according to the formula described by Yin et al. (2001). A set of 10-fold serial dilutions from 2 replicates was used to generate...

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Table 1. Primer and probe locations for detection of *Myxobolus cerebralis*. Tₘ: melting temperature; FAM: fluorescent reporter; NFG-MGB: non-fluorescent quencher attached to minor groove binding molecule

<table>
<thead>
<tr>
<th>Primers and probe</th>
<th>Sequence (5’–3’)</th>
<th>Sequence position</th>
<th>Tₘ (°C)</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Hsp 70</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Forward primer (414F)</td>
<td>CGAAGAACTCAATGGCTGATCTT</td>
<td>414–436</td>
<td>58.2</td>
</tr>
<tr>
<td>Reverse primer (471R)</td>
<td>TTGGTCAAATTGGATACGTAAGA</td>
<td>471–493</td>
<td>59.3</td>
</tr>
<tr>
<td>Probe (448P)</td>
<td>(FAM)-TCTTGACCCCGTTGAGAA-(NFG-MGB)</td>
<td>448–466</td>
<td>68.3</td>
</tr>
<tr>
<td><strong>18S rDNA</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Forward primer (811F)</td>
<td>TGAATAAATCAGAGTGCTAAAGC</td>
<td>811–833</td>
<td>56.6</td>
</tr>
<tr>
<td>Reverse primer (937R)</td>
<td>TTGGTGCTGTATGCTGTAACTG</td>
<td>937–958</td>
<td>55.3</td>
</tr>
<tr>
<td>Probe (888P)</td>
<td>(FAM)-TGTGACAAATAGCG-(NFG-MGB)</td>
<td>888–892</td>
<td>64.2</td>
</tr>
</tbody>
</table>
standard curves from the above plasmids for subsequent QPCR analysis. Linearity was determined through regression of the standard curve. Quantification of target DNA was determined by comparing critical threshold (Ct) values, which represents that portion of the exponential level of fluorescence that can be detected in the QPCR reaction from unknown samples with that derived from a known standard. Ct values are inversely proportional to the starting quantity of template DNA and can therefore be used as a relative indicator of template material in the absence of a standard curve. Prior to analysis, infected fish samples and plasmid standards were run in unison to ensure that amplification profiles were similar and that samples would fall within the range of the standard curve (data not shown).

**Assay specificity.** To demonstrate assay specificity, controls consisted of DNA isolated from several closely related organisms including *Myxobolus neurobious, M. insidiosus* and *M. squamalis*. In addition, DNA preparations from *Ceratomyxa shasta* and *Henneguya salmonicola*, 2 parasites known to infect salmonids (Noga 1996, Hoffman 1999), were also used to test assay specificity. We obtained 20 rainbow trout fry *Oncorhynchus mykiss* from a group of fish that had previously been determined negative for infection by nested PCR and histological analyses (Cavender et al. 2003), and used these to test for non-specific amplification of host tissue. These samples were randomly distributed throughout a 96-well optical plate and analyzed by PCR using both primer/probe sets.

**Assay sensitivity.** Triactinomyxon (TAM) spores obtained from C. Whipp at Oregon State University and E. MacConnell at Montana Fish, Wildlife and Parks, were enumerated using a glass microscope slide, 2 µl pipette and a light microscope. Nucleic acid was extracted from 20 TAM spores using standard reagents, microfiltration tubes and standard protocols outlined in Qiagen's DNeasy tissue kit. The nucleic acid concentration of this extraction was determined using a spectrophotometer. To demonstrate assay sensitivity, detection limits were examined with 10- and 2-fold serial dilutions from the above preparation using primer/probe combinations developed for both the Hsp 70 and 18S rDNA sequences. In addition, single TAM spores were placed in 5 µl of ddH$_2$O (double distilled water), denatured for 10 min at 100°C, and examined by QPCR analysis.

**Histopathology and PCR analysis.** Samples for use in this study were obtained from 107 rainbow trout fry exposed to *Myxobolus cerebralis* under natural river conditions (Cavender et al. 2003). Following exposure, fish were maintained until 900 CTUs (Celsius temperature units) had accumulated to allow pathology and early spore development to occur. At 900 CTUs, fish were euthanized with 200 mg 1$^{-1}$ tricaine methane sulfonate (MS-222; Argent Laboratories), decapitated, and their heads split sagitally. From each specimen, 1 half-head was processed for histological analysis at Washington’s Animal Disease and Diagnostic Laboratory (WADDL) located at Washington State University (Pullman). Slides were stained with hematoxylin and eosin, examined under a light microscope, and graded for infection intensity according to the MacConnell/Baldwin (M/B) scale of infection for *M. cerebralis* (Baldwin et al. 2000, as modified by Andree et al. 2002). Infection severity, as determined by histopathology, ranged from 0–5, with 0 indicating absence and 5 representing the most severe grade of infection.

The second half-head from each specimen was prepared for PCR analysis using a 6 mm biopsy punch (Miltex Instruments Company) to remove tissue containing cartilage and bone located at the base of the skull. Nucleic acid extractions and reaction conditions were performed as described by Cavender et al. (2003). All samples were analyzed with a single-round PCR analysis (Baldwin & Myklebust 2002). When examining lightly infected fish, it has been demonstrated that single-round PCR is less sensitive than the nested PCR for *Myxobolus cerebralis* (Kelley et al. 2004). Therefore, those samples determined negative by single-round PCR were further examined by nested PCR following previously established protocols (Andree et al. 1998). Tissues from fish examined by histopathology and PCR were analyzed using QPCR to provide a relative comparison of specificity and sensitivity between diagnostic methods.

**Statistical analysis.** To ensure that replicates for both Hsp 70 and 18S rDNA could be combined for statistical analysis, differences in copy number between duplicate wells were analyzed using a paired t-test. As no difference between replicate wells for either Hsp 70 (p = 0.94) or 18S rDNA (p = 0.82) was found, replicates were combined and mean values were used for further analysis. Copy numbers for both sequences did not meet normality assumptions and were subsequently log$_{10}$-transformed prior to examination. Detection prevalence for *Myxobolus cerebralis*-infected fish was analyzed as a categorical variable across all diagnostic methods using a chi-square test for proportion differences. Linearity for both standards was evaluated using regression of the linear curve as calculated by the iCycler real-time PCR detection system. Association between histology scores and mean copy number from the Hsp 70 and 18S rDNA sequences was examined using general linear regression. Significance throughout this study was defined as p < 0.01 and statistical analysis was conducted using the SAS software program, Version 8.2.
RESULTS

PCR amplification and species-specificity

Initial amplification of the appropriate nucleotide sequence for the Hsp 70 and 18S rDNA genes was done by PCR analysis using 150 and 20 ng µl⁻¹ of *Myxobolus cerebralis* DNA, respectively. Amplification of the 80 base-pair region from the Hsp 70 sequence and the 140 base-pair region from the 18S rDNA sequence was verified by agarose gel (2.0%) electrophoresis following PCR amplification (data not shown). Specificity of primer/probe combinations developed for both sequences was then optimized for QPCR analysis using DNA preparations from rainbow trout (host DNA only) and 5 fish parasites common to salmonids of the Pacific NW and capable of coexisting with *M. cerebralis* (Hoffman 1999). Both sets were determined to be specific only for *M. cerebralis* and did not react with host DNA (data not shown).

Assay sensitivity

A set of 10-fold serial dilutions of DNA extracted from 20 TAM spores (concentrations ranging from 25 to 0.25 pg µl⁻¹) was amplified using QPCR analysis. Positive amplification for both the Hsp 70 and 18S rDNA sequences was equivalent to 12.5 pg µl⁻¹ (Table 2). A set of 2-fold serial dilutions of DNA extracted from TAM spores (concentrations ranging from 25 to 1.6 pg µl⁻¹) was also amplified by QPCR. The lowest detectable DNA concentrations for the Hsp 70 and 18S rDNA sequences were equivalent to 12.5 and 6.3 pg µl⁻¹, respectively. Primer/probe combinations for both genes were capable of detecting single TAM spores following a short denaturing protocol. However, the copy number for single TAM spores ranged from 1–7.

Table 2. Results summarizing sensitivity of Hsp 70 and 18S rDNA QPCR analysis. Amplification above no template control (copy number = 0) was considered positive

<table>
<thead>
<tr>
<th>Triactinomyxon spores Dilution series</th>
<th>DNA conc. (pg µl⁻¹)</th>
<th>Hsp 70</th>
<th>18S rDNA</th>
</tr>
</thead>
<tbody>
<tr>
<td>20</td>
<td>1</td>
<td>25000</td>
<td>+</td>
</tr>
<tr>
<td>2</td>
<td>1/10</td>
<td>2500</td>
<td>+</td>
</tr>
<tr>
<td>0.2</td>
<td>1/10</td>
<td>250</td>
<td>+</td>
</tr>
<tr>
<td>0.02</td>
<td>1/10</td>
<td>25</td>
<td>+</td>
</tr>
<tr>
<td>0.01</td>
<td>1/1</td>
<td>12.5</td>
<td>+</td>
</tr>
<tr>
<td>0.005</td>
<td>1/1</td>
<td>6.3</td>
<td>–</td>
</tr>
<tr>
<td>0.0025</td>
<td>1/1</td>
<td>3.2</td>
<td>–</td>
</tr>
<tr>
<td>0.0013</td>
<td>1/1</td>
<td>1.6</td>
<td>–</td>
</tr>
</tbody>
</table>

Standard design and linearity

Nucleotide length and plasmid concentration was used to calculate copy number for each standard. Thus, 1 µg of DNA was calculated to be equivalent to 2.9 × 10⁹ and 2.27 × 10⁹ copies for the Hsp 70 and 18S rDNA sequences, respectively. We analyzed 10-fold serial dilutions of each standard by QPCR, and their Ct values were plotted against starting quantity for each plasmid. Regression of the slope for the Hsp 70 standard was −3.435 and the equation of the line was $y = -3.435x + 38.195$. Regression of the slope for the 18S rDNA standard was −3.461 and the equation of the line was $y = -3.461x + 35.018$. Linearity was demonstrated for both the Hsp 70 and 18S rDNA standards, with correlation coefficients corresponding to 0.995 and 0.997, respectively (Fig. 1).

Analysis of experimentally infected fish

Sentinel rainbow trout previously exposed to *Myxobolus cerebralis* under natural river conditions (Cavender et al. 2003) were examined for infection by histopathology, PCR, and QPCR analyses. Of the 107 fish examined, 32 were determined negative for infection by histology (Table 3). Fish analyzed by histology were also examined by single-round PCR, nested PCR, and QPCR using the primer/probe combinations developed for the Hsp 70 and 18S rDNA sequences. All fish that tested negative for infection by single-round PCR were also negative by nested PCR. Therefore, data from this analysis were combined and hereafter referred to as PCR analysis. Of 32 fish determined negative by histology, 9 (28%) were positive for infection by PCR and QPCR analyses. All fish determined positive for infection by histology were also positive by PCR and QPCR analyses. Detection by the PCR-based methodology was shown to be more sensitive, but when analyzed across all severity levels was not significantly greater than that observed for histology ($p = 0.213$). However, significantly greater detection of *M. cerebralis*-infected fish was shown to occur by PCR and QPCR among those fish determined negative by histology ($p = 0.008$).

Infection severity and genetic copy number

The copy number for the Hsp 70 gene increased with increasing histology scores of 0–4, but decreased with a histology score of 5 (Fig. 2). The copy number for the 18S...
rDNA gene increased with increasing histology scores of 0–3, but decreased with histology scores of 4–5. The observed decrease occurred gradually for the Hsp 70 gene and was abrupt for the 18S rDNA gene. The copy number for the Hsp 70 and 18S rDNA sequences ranged from 0–1880 and 0–11 200, respectively (Fig. 2). Significant correlation (p ≤ 0.01) between mean histology scores and mean genetic copy number did not occur when analyzed from 0–5 for either the Hsp 70 (R² = 0.74, p < 0.03) or the 18S rDNA (R² = 0.54, p < 0.09) sequences. However, the mean copy number did correlate significantly (R² = 0.96, p = 0.003) with mean histology scores of 0–4 for the Hsp 70 sequence. This correlation was not found between histology scores (0–4) and copy number for the 18S rDNA gene.

Table 3. *Oncorhynchus mykiss*. Histology, PCR and QPCR analysis of rainbow trout exposed to *Myxobolus cerebralis* under natural river conditions. Results from single and nested PCR analysis were combined and used to calculate PCR positive column. *Significant differences compared to histology

<table>
<thead>
<tr>
<th>Histology score</th>
<th>Positive by histology</th>
<th>PCR positive</th>
<th>QPCR Hsp 70</th>
<th>18S rDNA</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>0/32</td>
<td>9/32*</td>
<td>9/32*</td>
<td>9/32*</td>
</tr>
<tr>
<td>1</td>
<td>9/9</td>
<td>9/9</td>
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<td>9/9</td>
</tr>
<tr>
<td>2</td>
<td>10/10</td>
<td>10/10</td>
<td>10/10</td>
<td>10/10</td>
</tr>
<tr>
<td>4</td>
<td>33/33</td>
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<td>12/12</td>
<td>12/12</td>
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</tr>
</tbody>
</table>
DISCUSSION

Methods currently used to identify *Myxobolus cerebralis* and measure infection severity within infected fish can be time-consuming, labor-intensive and often subjective in their ability to distinguish the parasite from other myxozoan species. To improve diagnostic capabilities for this parasite, real-time QPCR assays targeting the Hsp 70 and 18S rDNA genes of *M. cerebralis* were developed. Sensitivity, specificity and potential correlation of assay results to infection severity (as measured by histology scores) were investigated using rainbow trout fry exposed to the parasite under natural river conditions.

In the present study, specificity of a QPCR analysis for *Myxobolus cerebralis* using 18S rDNA and Hsp 70 probe/primers was verified by testing cross-reactivity with DNA extracted from 20 uninected rainbow trout (negative controls) and 5 fish parasites. No amplified product was observed for host or non-*M. cerebralis* DNA, suggesting that both primer/probe combinations were specific for the target sequences. Assay sensitivity was further defined by comparing scores from 0–5 for either probe/primer set (Fig. 2). At higher histology scores, a drop in copy number was observed for the Hsp 70 and 18S rDNA sequences, the reason for such a decrease is unclear, but could be related to extraction inefficiencies at high parasite levels, the stimulation of a host immune response to infection, and/or inadvertent sampling bias (histology only provides a cross-sectional observation). Pathological conditions associated with increasing infection intensity by *M. cerebralis* include formation of myxospores and leukocyte infiltration around infected tissues (Andree et al. 2002). As myxospores begin to form, the ability to effectively extract nucleic acids decreases (Andree et al. 1997). Therefore, increasing parasite burden associated with elevated infection severity may have resulted in decreased extraction efficiency. The efficiency of extraction was consistent between different tubes. Although fish were sampled at 900 CTUs following exposure, spore development at high infection levels would be greater, and consequently the quantity or efficiency of DNA extraction may have been compromised due to the presence of mature spores. Alternatively, since increased histology scores can represent leukocyte infiltration associated with high infection levels (Andree et al. 2002), such a host response may reduce pre-spore developmental stages and spore numbers but fish may still exhibit high histology scores. A final possibility that might explain the decrease in copy number at high histology scores could relate to parasite distribution within the cranial region. This distribution probably varies and could create a sampling bias between that half of the head examined for histology and that half processed for PCR and QPCR analyses. Although a PTD analysis might have clarified some of these interpretations, this method could not be incorporated due to limited spore development during the period used to generate infected material and the sampling regime employed in this study.

As stated, a significant linear correlation between copy number and histology scores of 0–4 was identified for the Hsp 70 sequence. This observation indicates a potential for combining QPCR diagnostics with a measure of infection severity, but further testing is required to validate this. Until recently, the scale of infection used by many researchers and diagnostic laboratories was based on a scale of 0–4 (Hedrick et al. 1999, Baldwin et al. 2000) rather than 0–5. The previous scale may correlate better with QPCR results, but variability may still occur. Testing of fish infected with the parasite at various life stages is needed.

Overall, the parasite copy number obtained from the Hsp 70 sequence was consistently lower (0–1880) than that detected using the 18S rDNA sequence (0–11 200). It has been suggested that 104 copies of the 18S rDNA sequence are present in the genome of mature *Myxobolus cerebralis* spores (Kelley et al. 2004), but the copy number of the Hsp 70 gene is currently unknown. Our data suggest that the Hsp 70 gene may be found in lower numbers than the 18S rDNA gene. Previous studies have demonstrated that single TAM spores can contain 6 structural cells and up to 64 sporoplasms capable of infecting a susceptible host (El-Matbouli et al. 1995); therefore a minimum of
70 copies per spore could have been demonstrated from an individual TAM spore. Although both primer/probe sets developed for QPCR analysis were capable of detecting single TAM spores, copy number ranged between 1 and 7 per spore. The above discrepancy is probably a result of an inability to extract and capture total DNA from spores using the denaturing protocol employed.

This study has demonstrated that during early stages of infection (900 CTUs), QPCR is a sensitive detection method, and could potentially predict infection severity at histology scores of 4 and below using an Hsp 70 probe/primer set. Although the 18S rDNA PCR was more sensitive (6.3 pg µl⁻¹) when detecting the parasite than the Hsp 70 PCR (12.5 pg µl⁻¹) under the reaction conditions tested, significant correlation between copy number and histology scores up to 4 were only found for the Hsp 70 sequence. These data suggest that QPCR would be a useful tool for detecting *Myxobolus cerebralis* in infected samples and might provide a relative indication of infection severity. Advantages from a technique of this type include a high degree of sensitivity associated with PCR methodology, high sample throughput, and the ability to detect and quantify target sequences within a 1-step process. However, it should be noted that in situations where regulatory actions may result from diagnosis of *M. cerebralis*-infected fish, all molecular assays should be coupled with traditional diagnostic techniques to avoid misdiagnosis.

The results of this study demonstrate the potential for QPCR to be used as a detection method for *Myxobolus cerebralis*. Future testing should include additional fish pathogens that may co-occur with *M. cerebralis* and controlled laboratory experiments with fish exposed to known spore concentrations. Fish should be sampled over the course of an infection to better identify relationships between copy number and infection severity. In addition, examining DNA from mature myxospores, infection within the secondary host (*Tubifex tubifex*), and testing of water samples would further validate the range of applications for this technique.

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