Development and application of a real-time quantitative PCR assay for determining CYP1A transcripts in three genera of salmonids

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Abstract

The expression of CYP1A (cytochrome P450 1A) can be induced by a large array of aromatic and organic compounds in teleost fishes. We developed a real-time quantitative PCR assay useful for measuring β-naphthoflavone (BNF) induction of liver CYP1A mRNA in four salmonid species. First, to obtain necessary information for the design of a cRNA standard, full-length CYP1A cDNA sequences were determined for two Salvelinus species, lake trout (S. namaycush) and brook trout (S. fontinalis). Each cDNA was found to share the same characteristics with known CYP1A sequences of Atlantic salmon (Salmo salar) and rainbow trout (Oncorhynchus mykiss): a start codon, conserved heme-binding region, putative poly-adenylation signal, stop codon, relatively long 3′-untranslated region (UTR; >1 kb), and a protein length of 523 amino acid residues. The brook trout and lake trout CYP1A cDNA’s were 2636 and 2672 base pairs (bp) in length and shared greater than 97% coding region sequence identity with Atlantic salmon and rainbow trout CYP1A’s. Next, using the generated sequence information, we developed a CYP1A-specific real-time quantitative PCR assay. Primers and a fluorescent-labeled probe were designed from a 68 bp region that was found to be conserved among salmonid CYP1A genes. The assay was designed to allow for simultaneous comparison of CYP1A expression among each experimental group. Finally, groups (n = 4–8) of hatchery-raised Atlantic salmon, brook trout, lake trout, and rainbow trout were given an intraperitoneal injection of a corn oil control, 25 mg kg⁻¹ BNF, or 50 mg kg⁻¹ BNF and sacrificed after 48 h. Liver tissue was collected and CYP1A mRNA levels were estimated. In all species, BNF treated fish showed 1.8–3.0 orders of magnitude higher CYP1A than control fish. The CYP1A induction levels were not different in fish treated with both dosages. Mean base levels of CYP1A expression ranged from 7.24 × 10⁶ (rainbow trout) to 1.30 × 10⁷ (brook trout) transcripts g⁻¹ total RNA. Mean induced levels of CYP1A expression ranged from 1.07 × 10⁸ (lake trout) to 1.30 × 10⁹ (brook trout) transcripts g⁻¹ total RNA.

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1. Introduction

Cytochrome P4501A’s (CYP1A) constitute a ubiquitous family of proteins associated with the detoxification of organic compounds such as PCB (polychlorinated biphenyl), PAH (polyaromatic hydrocarbons), and dioxin (Mansuy, 1998; Nelson et al., 1996; Kleinow et al., 1987). These compounds are documented to induce the CYP1A gene in a variety of tissues of many fish species (Levine and Oris, 1999; Hahn et al., 1998; Gooneratne et al., 1997; Nelson et al., 1996; Smolowitz et al., 1992; Müller et al.,...
1989). Consequently, changes in CYP1A gene expression have been used as a biomarker for contaminant exposure in fish populations (Cousinou et al., 2000; Miller et al., 1999; Campbell and Devlin, 1996; Levine et al., 1994; Stegeman et al., 1992).

A variety of techniques have been applied to estimate CYP1A induction in fish. Protein levels can be measured by determining EROD (ethoxyresorufin-O-deethylase) activity (Schlezinger and Stegeman, 2001; Andersson et al., 1985; James and Bend, 1980), immunohistochemistry (Stegeman et al., 2001; Van Veld et al., 1992; Smolowitz et al., 1991), enzyme-linked immunosorbent assay (Sarasquete and Segner, 2000), and Western blotting (Grøsvik et al., 1997). Recently, CYP1A gene expression has been estimated from mRNA levels through Northern blotting (Grøsvik et al., 1997), slot-blotting, or quantitative PCR (Rees et al., 2003; Miller et al., 1999; Campbell and Devlin, 1996). Of all of these methods, quantitative PCR appears to be the most sensitive (Vanden Heuvel et al., 1993). It has been used to assess impact of environmental pollution in marine ecosystems using emerald rockcod (Trematomus bernacchi, Miller et al., 1999), guilthead seabream (Sparus aurata, Cousinou et al., 2000), and grey mullet (Liza aurata, Cousinou et al., 2000). Likewise, numerous quantitative PCR assays have been developed to study CYP1A induction in freshwater teleosts such as Pacific salmon (Oncorhynchus tshawytscha, Campbell and Devlin, 1996) and Atlantic salmon (Salmo salar, Rees et al., 2003).

The goal of this study was to develop a single real-time quantitative PCR assay and use it to estimate CYP1A levels in at least three genera of salmonids: Oncorhynchus, Salmo, and Salvelinus. This is feasible because orthologous CYP1A genes are often highly conserved. Coding sequences of the CYP1A gene in Atlantic salmon (Rees et al., in press) and rainbow trout (Berndtson and Chen, 1994; Heilmann et al., 1988), for instance, share more than 96% of sequence identity. However, CYP1A sequences from Salvelinus species are not available. Therefore, our first objective was to clone CYP1A in two Salvelinus species, lake trout (S. namaycush) and brook trout (S. fontinalis). Our second objective was to develop a quantitative PCR assay useful for all three genera of salmonids. The third objective was to use the anticipated assay to determine and compare the effect of β-naphthoflavone (BNF) treatment on liver CYP1A levels in all species of the three genera.

2. Materials and methods

2.1. Animals, BNF induction, and acquisition of tissues

All species of salmonids used in this study were acquired from nearby fish hatcheries. Juvenile lake trout and brook trout (11 g ± 2 g mean weight, 11 cm ± 2 cm mean length) were acquired from Marquette State Fish Hatchery (Marquette, MI, USA), juvenile Atlantic salmon (20 g ± 3 g mean weight, 13 cm ± 2 cm mean length) were collected from Lake Superior State University Fish Hatchery (Sault Ste. Marie, MI, USA), while juvenile rainbow trout (11 g ± 2 g mean weight, 11 cm ± 2 cm mean length) were collected from Wolf Lake State Fish Hatchery (Mattawan, MI, USA). All fish were held at Michigan State University where they were acclimated for two weeks at 12 °C in 800-l flow-through tanks (well water, 600 l h⁻¹). A 12 h light-dark cycle was maintained during the acclimation and experiment period. Fish were fed Purina AquaMax© Grower 400 (lot A-5D04; Purina Mills, Inc.; St. Louis, MO) daily at a level of 1.5% body weight. Two days prior to injection, fish were taken off of feed. Individuals were randomly sampled and given an intraperitoneal injection of a corn oil control or β-naphthoflavone (BNF, Sigma Chemical Corp.; St. Louis, MO) dissolved in corn oil at doses of either 25 or 50 mg kg⁻¹ body weight. Fish were then placed in a 40-l flow-through aquarium (20 l h⁻¹) for 48 h and then sacrificed using an overdose of MS-222 (Sigma Chemical Corp.). BNF induction of CYP1A reaches maximum in 48 h (Grøsvik et al., 1997). Tissues (gill, liver, and brain) were immediately collected and stored in RNALater® at −80 °C (Ambion; Austin, TX). In addition, gill and liver tissue from one induced lake trout and one induced brook trout were collected and used for cloning of the CYP1A gene in each species.

2.2. RNA isolation

RNALater® preserved tissues were homogenized and extracted for total RNA isolation using Trizol...
Reagent (Life Technologies; Carlsbad, CA) according to the manufacturer’s protocol. RNA samples were incubated at 37°C with RNase-free DNase I (Roche Molecular Biochemicals; Mannheim, Germany) then re-suspended in 20–50 μl of diethylpyrocarbonate-treated water (DEPC-H2O) and quantified (Sambrook et al., 1989) using a GeneQuant pro RNA/DNA calculator (Amersham Biosciences; Piscataway, NJ). For long-term storage, RNA samples were supplemented with three volumes of 95% ethanol, 1/10 volume of 3 M sodium acetate, and stored at −80°C (Sambrook et al., 1989).

2.3. Cloning of full-length cDNA’s encoding lake trout and brook trout CYP1A genes

We followed the strategy and procedures of Rees et al. (in press) for cloning these two CYP1A genes. Briefly, RACE was carried out using the Advantage II RACE system (Clontech; Palo Alto, CA) according to the manufacturer’s protocol. One μg of total RNA from lake trout gill and brook trout liver was used as a template for synthesis of 3′-RACE Ready cDNA. We conducted a 3′-RACE with a gene specific primer (WML56 5′-CGG CTC A TT TGG CTC A TA ACG GAA GA T-3′) designed from the 5′-UTR sequence of Atlantic salmon CYP1A (Rees et al., in press, GenBank accession number AF361643). This RACE insured that all functional domains including the 5′-UTR and entire 3′-UTR would be included in the cloned cDNA. After cloning, both RACE products were sequenced by the Plant Biology DNA Sequencing Facility, Michigan State University.

2.4. Phylogenetic analysis

The coding region of brook trout, lake trout, and Atlantic salmon CYP1A was aligned to the coding region of a sample of P450 genes using the CLUSTAL W algorithm. Genetic relationships and distances were generated using the Neighbor-joining method. Genes selected for this analysis comprised teleost CYP genes representing families 1–4 (refer to Table 1 for GenBank accession numbers).

2.5. In vitro transcription of cRNA standard

Separate plasmids containing either the full CYP1A cDNA sequence from lake trout, brook trout, and Atlantic salmon (Rees et al., in press) were obtained by standard cloning procedures and sequenced as described previously. To design a cRNA standard, a 491 bp conserved region of the CYP1A gene was amplified from the Atlantic salmon CYP1A clone using the following primers and conditions: forward primer WML 169 5′-TAA TAC GAC TCA CTA TAG GCT GTC TTG GGC TGT TGT GTA CCT TGT G-3′, reverse primer WML 170 5′-TTT TTT TTT TTT TTT GGA GCA GGA TGG CCA AGA AGA GGT AG-3′, 1 cycle at 94°C for 4 min, 40 cycles 94°C for 5 s and 72°C for 2 min, and 1 cycle at 72°C for 5 min as added extension. The PCR product contained a

Table 1

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<tr>
<th>Name in analysis (common name)</th>
<th>Reference</th>
<th>GenBank accession number</th>
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<tr>
<td>Anguilla japonica CYP1A (Japanese eel)</td>
<td>Minato et al. (1999) (unpublished)</td>
<td>AB020414</td>
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<tr>
<td>Danio rerio CYP1A (zebrafish)</td>
<td>Yamazaki et al. (2002) (unpublished)</td>
<td>AB078927</td>
</tr>
<tr>
<td>Oncorhyncus mykiss CYP1A (rainbow trout)</td>
<td>Berndtson and Chen (1994)</td>
<td>S69278</td>
</tr>
<tr>
<td>Oncorhyncus mykiss CYP1A3</td>
<td>Berndtson and Chen (1994)</td>
<td>S69277</td>
</tr>
<tr>
<td>Salmo salar CYP1A (Atlantic salmon)</td>
<td>Rees et al. (2003)</td>
<td>AF361643</td>
</tr>
<tr>
<td>Salvelinus namaycush CYP1A (lake trout)</td>
<td>Rees and Li, (this paper)</td>
<td>AF539414</td>
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<tr>
<td>Salvelinus namaycush CYP1A27</td>
<td>Rees and Li, (this paper)</td>
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<td>Oryzias latipes CYP3A (Japanese medaka)</td>
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<tr>
<td>Dicentrarchus labrax CYP4D11 (sea bass)</td>
<td>Salvarani et al. (1998)</td>
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5'-T7 promoter, 454 bp of CYP1A sequence including the region of the real-time amplicon, and a poly dT tail at the 3'-end. This product was then diluted 1/100 with deionized water, re-amplified and scaled up with the same reaction conditions. Additional handling, amplification, and purification of the cRNA standard were performed as described previously (Rees et al., 2003).

2.6. Quantitative PCR primer and probe design

PCR primers and the fluorescent-labeled probe were designed to conform to several criteria. First, primers should only amplify the CYP1 family of P450 genes and not any other P450 family (i.e. CYP2 or CYP3). In addition, they should anneal to an existing region on the CYP1A gene that was highly conserved (>99%) over all the known salmonid CYP1A sequences. These criteria were met by performing two separate multiple sequence alignments (CLUSTAL W algorithm, DNASTAR; Madison, WI) with known teleost CYP genes. The first alignment was performed with Atlantic salmon CYP1A against six different rainbow trout CYP genes representing families 1–3. Then, a second alignment was performed with all of the known salmonid CYP1A sequences from three different salmonid genera: Oncorhynchus, Salmo, and Salvelinus. We selected a 68 bp region that was conserved among CYP1A genes of Oncorhynchus, Salmo, and Salvelinus and was different from other families of rainbow trout CYP genes.

2.7. RT-PCR

Reverse transcription was performed as described previously (Rees et al., 2003). Each PCR reaction consisted of 25 µl of 2× TaqMan® Universal PCR master mix (Applied Biosystems; Branchburg, NJ), 300 nM of each primer, 100 nM of the TaqMan® probe (5'-6-FAM, 3'-TAMRA quencher), 1 µl of cDNA template, and DI water to a final volume of 50 µl. Reactions were then analyzed on an ABI 7700 real-time PCR thermalcycler (Applied Biosystems) under the following conditions: 50°C for 2 min, 95°C for 10 min, and 40 cycles of 95°C for 15 s followed by 60°C for 1 min. Amplification plots were generated and CYP1A mRNA levels were estimated against a standard curve.

2.8. Primer and probe optimization

Real-time PCR primers (forward WML158 5'-CCA ACT TAC CTC TGC TGG AAG C-3' and reverse WML159 5'-GTT GAA CGG CAG GGA GGA-3') were optimized for quantitative PCR by performing PCR reactions with nine separate concentration combinations in quadruplicate and determining which combination produced the largest ∆Rn. ∆Rn (normalized reporter) represents the signal to noise ratio and indicates the magnitude of the signal generated by a given set of PCR conditions for more information, consult the Applied Biosystems TaqMan® Universal PCR Master Mix Protocol. Once the primer concentration was chosen, an additional set of reactions was set up to optimize the probe (WML160 5'-TTC A TC CTG GAG A TC TTC CGG CAC TC-3') concentration for the chosen primer concentration. Five separate probe concentration reactions were used in quadruplicate and analyzed to see which concentration produced the smallest Ct (threshold cycle). Ct values represent the cycle of amplification at which a PCR reaction reaches a statistically significant increase in ∆Rn (consult the Applied Biosystems TaqMan® Universal PCR Master Mix Protocol for more information). The lowest Ct value in these optimization reactions indicates the concentration at which optimal probe binding occurs and point of highest sensitivity for detecting specific template. For all of the above reactions, Atlantic salmon liver cDNA was used as the PCR template.

2.9. Standard curve

A standard curve for each set of samples was generated by performing RT-PCR on a dilution series of the recombinant cRNA standard. The concentration of the standard molecule was estimated in terms of molecules. A 10× dilution series was carried out from 10^5 to 10^10 molecules. Amplification plots were analyzed on the ABI 7700 and Ct values for each of the reactions in the dilution series were calculated. Ct values were plotted against starting quantity of RNA template to generate the standard curve (refer to Fig. 5 for a representative standard curve). A standard curve was generated for each plate analyzed.
2.10. Statistical analysis

Transcript numbers of CYP1A µg⁻¹ total RNA were calculated from the appropriate standard curve and log transformed. Data were analyzed using a two-way analysis of variance (ANOVA; Statistical Analysis Systems v.8, Cary, NC). All pairwise comparisons were tested for significance by using a Tukey–Kramer adjustment (Kramer, 1956).

3. Results

3.1. Brook trout and lake trout CYP1A sequences

3′-Long distance RACE on brook trout cDNA produced a PCR fragment of 2636 bp (Fig. 1). Sequence analysis indicated this product included 33 bp of the 5′-untranslated region (UTR), a 1569 bp coding region, and a 1034 bp 3′-UTR containing three AUUUA sequences. It encodes a protein of 523 amino acid residues. The sequence also possessed all major functional domains and characteristics of previously discovered CYP1A molecules including the heme-binding cysteine (position 463), arginine codon (position 246) integral to enzymatic function, stop codon (position 523), and poly-adenylation signal.

A lake trout CYP1A fragment of 2672 bp long was also cloned and sequenced (Fig. 2). This clone contained 28 bp of the 5′ UTR, a 1569 bp coding region, and a 3′-UTR of 1073 bp in length containing two AUUUA sequences. It also encodes a protein of 523 amino acid residues and contains all of the major functional domains typical of CYP1A’s as described above.

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**Fig. 1.** cDNA and deduced amino acid residue sequence of brook trout CYP1A (GenBank accession number AF539414). The start codon, arginine residue critical to enzymatic function (position 246), heme-binding cysteine codon (position 463), stop codon (position 523), AATTA (AUUUA) sequences, and putative poly-adenylation signal are all underlined.

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Fig. 2. cDNA and deduced amino acid residue sequence of lake trout CYP1A (GenBank accession number AF539415). The start codon, arginine residue critical to enzymatic function (position 246), heme-binding cysteine codon (position 463), stop codon (position 523), ATTTA (AUUUA) sequences, and putative polyadenylation signal are all underlined and boldfaced.

3.2. Phylogenetic analysis

For the coding region, the brook trout and lake trout CYP1A genes described here share ~97% sequence identity with each other. These two genes also share >97% sequence identity with Atlantic salmon and rainbow trout CYP1A genes. In addition, brook trout and lake trout CYP1A genes share between 70–80% nucleotide homology with other teleost CYP1A genes.

Multiple sequence alignment using the CLUSTAL W algorithm followed by construction of a phylogenetic tree using the Neighbor-joining method suggested that all of the salmonid CYP1A genes are highly related (Fig. 3). Minor differences in sequence data grouped Atlantic salmon and rainbow trout CYP1A genes together. Lake trout CYP1A was genetically closest to rainbow trout CYP1A3. Gene names and GenBank accession numbers used in the phylogenetic analysis can be found in Table 1.

3.3. Primer and probe design and optimization

As discovered from multiple sequence alignments, we determined that the primers and probe chosen would be sufficient to amplify CYP1A from each of the salmonid species listed without cross amplification of alternative teleost CYP genes. The region chosen represents an amplicon of 68 bp long at nucleotides 1103–1170 of the coding region of Atlantic salmon CYP1A. Refer to Fig. 4 for a comparison of this region between salmonid species. The highest mean ΔRn (2.88) was found with both forward and reverse primers (WML158 and WML159) at 300 nM. Under these primer conditions the probe was found to produce the lowest mean Ct (22.96) at 100 nM.
Fig. 3. Phylogenetic analysis of cytochrome P450 genes. Multiple sequence alignment was carried out using the Clustal W algorithm (only coding regions of each respective gene were used). The phylogenetic tree and genetic distances were determined using the Neighbor-joining method.

3.4. Standard curve

The reactions for the standard curve were run on the same plate as all analyzed samples. Ct values were plotted against concentrations of cRNA standard transcripts and analyzed using linear regression. The standard curve had a slope of −3.8 and a coefficient of determination of 0.98 (Fig. 5). All Ct values of RNA extracted from each individual fell within the linear range of the standard curve.

3.5. CYP1A induction in liver of brook trout, lake trout, rainbow trout, and Atlantic salmon

CYP1A was induced approximately 100–1000 fold in all species injected with both dosages (25 mg kg⁻¹ body weight and 50 mg kg⁻¹ body weight; Fig. 6). A significant interaction was found between species and treatment making main effects irrelevant. Simple effects were determined for each factor by using the SLICE procedure (Statistical Analysis Systems...
Fig. 5. Standard curve for the real-time CYP1A quantitative PCR assay. A 10-fold dilution series was carried out for the cRNA standard from $10^2$ to $10^3$ molecules and amplified for 40 cycles during PCR. $C_t$ (cycle threshold indicating the first detection of CYP1A PCR product) values were plotted against initial concentration followed by standard linear regression ($r^2 = 0.98$). There was no statistical difference between BNF induction at 25 or 50 mg kg$^{-1}$ body weight in each species ($P > 0.45$). However, ANOVA analysis showed a difference between CYP1A mRNA levels of control and induced groups in each species ($P < 0.0001$). On average, all species had approximately $1 \times 10^6$ CYP1A transcripts mg$^{-1}$ total RNA at control levels and $1 \times 10^9$ CYP1A transcripts mg$^{-1}$ total RNA under induced conditions. Brook trout demonstrated higher mean basal levels of CYP1A than all other species (ANOVA $P < 0.01$) at $1 \times 10^6$ CYP1A transcripts mg$^{-1}$ total RNA. Among BNF induced fish, lake trout had lower levels of CYP1A transcripts at approximately $1 \times 10^8$ CYP1A transcripts mg$^{-1}$ total RNA (ANOVA $P < 0.02$) than all other treatment groups except Atlantic salmon treated with 50 mg kg$^{-1}$ BNF.

4. Discussion

It is evident that both of the PCR fragments cloned from lake trout and brook trout represent full-length cDNA clones of CYP1A genes. Each cDNA sequence has characteristics of a full-length teleost CYP1A cDNA: a start codon and a stop codon followed by a poly A tail, a heme-binding domain, an arginine codon integral to enzymatic function, and a rather large 3'UTR containing two (lake trout) and three (brook trout) AUUUA sequences. The coding region (1569 bp), which encodes a protein of 523 amino acid residues, is the same size as the rainbow trout and Atlantic salmon P450 1A protein. In addition, the brook trout and lake trout CYP1A genes show 97.9 and 97.6% sequence identity to Atlantic salmon CYP1A. The coding regions of CYP1A genes isolated to date in salmonids (Atlantic salmon, rainbow trout, Pacific salmon, brook trout, and lake trout) differ by no more than 3.9%. This high level of sequence identity confirms that the CYP1A gene is highly conserved and thus suitable for developing a real-time quantitative PCR assay to study CYP1A expression dynamics across several genera in response to contaminant exposure.

Real-time PCR analysis indicates that CYP1A induction in liver tissue of lake trout, brook trout, rainbow trout, and Atlantic salmon followed a consistent pattern. In all species, CYP1A expression was induced by BNF injection from approximately 1.8–3.0 orders of magnitude representing a 60–1000-fold difference in CYP1A levels between control and induced levels. This trend of induction was seen in fish injected with 25 mg kg$^{-1}$ and also 50 mg kg$^{-1}$ BNF. Previous quantitative PCR studies have also found a similar level of induction by BNF in Atlantic salmon (Rees et al., 2003) and Pacific salmon (Campbell and Devlin, 1996) in a variety of tissues (liver, kidney, gill, brain, and gonad). Absolute levels of CYP1A expression ranged from a low of approximately $5 \times 10^5$ molecules CYP1A $\mu$g$^{-1}$ total RNA in rainbow trout liver (control group) to a high of approximately $1 \times 10^9$ molecules CYP1A $\mu$g$^{-1}$ total RNA in rainbow trout liver (induced). The CYP1A levels reported here closely resemble CYP1A mRNA expression levels identified in a 28 day BNF induction time course in Pacific salmon liver tissue. Campbell and Devlin (1996) report that at time zero Pacific salmon liver shows similar levels of CYP1A at $5.00 \times 10^3$ molecules $\mu$g$^{-1}$ total RNA. After 28 days, this expression jumps 160-fold to $8.04 \times 10^7$ transcripts $\mu$g$^{-1}$ total RNA. However, the results from each of these experiments are 2–3 orders of magnitude lower than those determined using a competitive quantitative PCR (Rees et al., 2003). This
Fig. 6. Real-time PCR analysis of liver CYP1A levels in representative salmonid species (AS: Atlantic salmon, BT: brook trout, LT: lake trout, RBT: rainbow trout). Fish were administered an injection of corn oil only (control), 25 mg kg\(^{-1}\) body weight BNF (β-naphthoflavone), or 50 mg kg\(^{-1}\) body weight BNF. Total RNA (100 ng) was reverse-transcribed and amplified in real-time from each treatment group (\(n = 4–8\); sample size is indicated for each treatment group below each bar) after which CYP1A levels were estimated. Bars represent mean logarithmic values of CYP1A expression total RNA ± S.E.M. for each treatment group. Comparisons were made between induced and control levels using a Tukey–Kramer adjustment for multiple comparisons. Symbol (*) notes significance of induced groups over each respective control group at \(P < 0.0001\). Symbol (**) notes significantly higher CYP1A levels over other control groups at \(P < 0.01\).

The real-time PCR developed in this study is highly sensitive and versatile. Based on our standard curves, it can measure CYP1A mRNA expression levels down to \(\sim 1000\) transcripts of CYP1A µg\(^{-1}\) total RNA because the standard curve obtained in this assay covers a large linear range (\(10^{10}–10^3\) molecules) and allows for versatility in measuring CYP1A gene expression through a wide degree of environmental and laboratory conditions. At \(\sim 100\) transcripts, a strong signal was observed but the \(C_t\) fell out of the linear range of the standard curve. This high level of sensitivity and wide range of applicability will likely enable accurate measurement of CYP1A levels in wild fish from both pristine and highly polluted environments. Because many salmonid species are at threatened status or worse (U.S. Department of Interior, 2000, 2002), this quantitative PCR assay will complement the development
of a non-lethal gill biopsy method to monitor contaminant exposure in salmonid populations (Rees et al., in review) without sacrificing individual fish captured in the wild.

Furthermore, this assay makes measuring CYP1A gene expression among various species more accurate, comparable, and quicker because the same primers and probe are used for each species. This allows RNA from various tissues of multiple species to be analyzed on the same plate and compared to the same standard curve. This assay also minimizes the possibility of quantifying false positives such as non-specific PCR products because the probe is single-stranded and only binds to the target sequence. Fluorescence is not emitted unless this binding occurs; therefore, only fluorescence from specific binding is measured (Giuletti et al., 2001). In addition, the time-intensive process during the generation of a “pure” cRNA standard is minimized because the same standard can be used for multiple species. This type of application has been used in the recent past to detect and quantify the same infectious heamatopoietic necrosis virus (iHNV) in multiple salmonid species (Overturf et al., 2001). However, our study documented a use of the real-time PCR assay to measure expression levels of an orthologous gene across several members in a single teleost family.

From an environmental assessment standpoint, it is important to realize the benefits and limits of this quantitative PCR assay. Because the amplification conditions are optimized and primer and probe sequences are known, it will be possible for other laboratories to adopt this method in a matter of days. This approach has the advantage of providing rapid and accurate measures of CYP1A transcripts in various tissues. To develop a comprehensive understanding of CYP1A induction in fish responding to contaminant exposure, this method should be combined with methods that provide better spatial resolution, such as in situ hybridization and immunocytochemistry, and that provide estimated CYP1A protein levels, such as ELISA, Western blotting and EROD analysis.

In conclusion, we have developed a real-time quantitative PCR assay for analysis of CYP1A expression across three salmonid genera, Salmo, Oncorhynchus, and Salvelinus. In development of this assay, we confirmed that CYP1A genes across the salmonid family carry a high degree of sequence homology and is highly induced in liver tissue of BNF-exposed lake trout, brook trout, Atlantic salmon, and rainbow trout after 2 days. We also discovered some species-specific characteristics of CYP1A induction. Brook trout showed higher basal levels and induced levels than all other species. Lake trout showed the lowest induced levels of CYP1A expression. It is notable that brook trout and lake trout, which showed a significant difference in induction, were reared in the same hatchery since embryonic stage. The ecological and physiological implication of this species difference, however, will require further studies to clarify. Finally, the real-time assay has a high degree of specificity for generated CYP1A PCR products as well as a high degree of sensitivity detecting down to 1000 molecules CYP1A μl−1 total RNA.

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References


