PRESENCE OF ECOTOXICOLOGICALLY RELEVANT Pgp AND MRP TRANSCRIPTS AND PROTEINS IN CYPRINID FISH

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One of the most intriguing defence strategies which aquatic organisms developed through evolution is multixenobiotic resistance (MXR). The key mediators of MXR activity are ATP-binding cassette (ABC) transport proteins. They provide resistance of aquatic organisms by binding xenobiotics and extruding them from cells in an energy-dependent manner. Since Cyprinid fish species are common target in freshwater biomonitoring programs, we have studied the presence of two main MDR/MXR efflux transporters P-glycoprotein (Pgp, Abcb1) and MRP-like protein(s) (Abcc) in the liver of five Cyprinid species: common carp, European chub, sneep, barbel, and silver prussian carp. Their presence was evaluated on the mRNA and protein level. Various pairs of primers were designed to clone homologous fragments of MXR-related genes. At the protein level, we used Western blotting with specific monoclonal antibodies against human Pgp (Abcb1, Ab C219), MRP1 (Abcc1; Ab MRPm6) or MRP2 (Abcc2; Ab M2I-4). Transcripts of both key types of MXR transporters were identified in all species examined and here we provide the phylogenetic analysis of new partial sequences. Immunochemical determinations with mammalian antibodies failed to identify the presence of MRP(s), but Pgp expression was found in all five Cyprinid species. These results support that MXR is a defence system mediated by both Pgp and MRP types of ABC transport proteins.

KEY WORDS: ABC proteins, aquatic organisms, multixenobiotic resistance, MXR, phylogenetic analysis, Western blot analysis
chemicals of both natural and anthropogenic origin, commonly called MXR inhibitors or chemosensitisers (10,11). However, the recent discovery of non-Pgp efflux transporters in various mammalian tissues implies that MXR in aquatic biota may also be a multi-transporter mechanism. Among non-Pgp ABC proteins, members of a multidrug resistance-associated protein (MRP) subfamily ABCC have been shown to be toxicologically relevant (12-14). Our earlier studies (15-17) and studies of other groups recently demonstrated the expression of MRP-related genes in a number of fish and invertebrate species (18-20).

The primary goal of this study was to determine Pgp (Abcb1) and Mrp (Abcc) related mRNA transcripts in a group of freshwater cyprinidae, as follows: common carp (Cyprinus carpio), European chub (Squalius cephalus), sneep (Chondrostoma nasus), barbel (Barbus barbus), and silver Prussian carp (Carassius auratus gibelio). These species are typical representatives of the fish communities of continental European waters and are frequently used in biomonitoring programmes/studies (21, 22). Additionally, we tried to detect Pgp and MRPs on the protein level using commercially available antibodies directed against human Pgp and MRPs.

Identified sequences share a high degree of homology with the P-glycoprotein (Pgp, Abcb1), Mrp1 (Abcc1), Mrp2 (Abcc2), and Mrp3 (Abcc3) proteins, while phylogenetic analysis shows grouping of identified sequences within ABCB and ABCC subfamily of other organisms.

MATERIALS AND METHODS

Animals

Chub, sneep, barbel, and silver Prussian carp specimens were collected by electrofishing from the Sava River (Croatia), while common carp was provided by a local fish farm.

Detection of Pgp and Mrp related mRNA transcripts in fish liver

Primer pairs (Table 1) were designed from highly conserved regions of mammalian and fish ABCB1 and ABCC1/2 genes and obtained from Operon Biotechnologies (Köln, Germany) or Invitrogen (Carlsbad, CA, USA).

Total RNA was extracted from liver tissue using the Agilent Total RNA Isolation Mini Kit (Agilent Technologies, Palo Alto, CA, USA). The quality of RNA samples was analysed using an RNA 6000 Nano LabChip Kit (Agilent Technologies). Three micrograms of total RNA were then reverse transcribed using M-MuLV reverse transcriptase according to manufacturer’s instructions (RevertAid First Strand cDNA synthesis Kit, Fermentas, Ontario, Canada). Polymerase chain reaction (PCR) was performed in Biometra thermal Cycler (Goettingen, Germany) with 3 µL of cDNA in a total volume of 50 µL. The cycling parameters were as follows: denaturation at 94 °C for 2 min, 35 cycles of heat denaturation at 94 °C for 30 s, annealing at 54 °C for 30 s, polymerisation at 72 °C for

Table 1  Primers used for PCR amplification of cDNA corresponding to Pgp (Abcb1) and Mrp (Abcc)-related genes

<table>
<thead>
<tr>
<th>PRIMER PAIR</th>
<th>5’→3’</th>
<th>FISH</th>
<th>TARGETED GENE</th>
<th>IDENTIF. GENE</th>
</tr>
</thead>
<tbody>
<tr>
<td>A/B F/T</td>
<td>GCG GCT GTG GGA AGA GCA C/TGT TGT CTC CGT AGG CAA TGT T</td>
<td>All fishes</td>
<td>Pgp (Abcb1)</td>
<td>Abcb1</td>
</tr>
<tr>
<td>C/D F/R</td>
<td>ACCGAGATCGGAGAGAGG CTG TCC AGG ATG GTG TTG A</td>
<td>Chub</td>
<td>Prussian carp</td>
<td>Abcc2</td>
</tr>
<tr>
<td>E/F F/R</td>
<td>GGTGGAGGCTGCTGCTCT CTG TCC AGG ATG GTG TTG A</td>
<td>Common carp Barbel</td>
<td>Mrp1/2 (Abcc1/2)</td>
<td>Abcc3</td>
</tr>
<tr>
<td>G/H F/R</td>
<td>AAT GAC ACC ATA CAG GAC AAC ATC CA TCC ACT GCA GAC AGG GGG GC</td>
<td>Sneep</td>
<td></td>
<td>Abcc2</td>
</tr>
</tbody>
</table>
30 s, and final extension at 72°C for 7 min. Aliquots of each reaction were resolved by electrophoresis on 1.2% agarose gel in the TAE buffer, the gels were stained with ethidium bromide and PCR products visualised under UV light. The expected amplimers obtained from all cyprinid species were excised and eluted using MiniElute PCR Purification Kit (Qiagen, Hilden, Germany). Isolated cDNAs of less than 1,000 bp size were sequenced directly, while longer cDNAs were subcloned into a suitable plasmid vector and transformed into *Escherichia coli* (PCR Cloning Kit, Qiagen). Plasmids were isolated from overnight cultures of transformants using QIAprep Miniprep (Qiagen). cDNAs were sequenced on both strands by VBC-Genomics (Vienna, Austria).

**Fish tissue collection and Western blot analysis**

Within three hours after catch, fish were killed and livers dissected and stored in liquid nitrogen. Membrane vesicles were prepared as described by Cornwell et al. (23). Phenylmethanesulfonylfluoride (PMSF) serine protease inhibitor was substituted with a protease inhibitor cocktail (Sigma, St Louis, MO, USA). Membrane vesicle proteins were resolved on a 7.5% acrylamide gel at a constant voltage of 200 V in BioRad Mini-PROTEAN 3 Cell system; 25 mmol L⁻¹ Tris-HCl, pH 8.3, 192 mmol L⁻¹ glycine, and 0.1% SDS running buffer. Electrophoretic transfer to Immobilon-P membrane (Millipore, Schwalbach, Germany) in BioRad Mini Trans-Blot followed the manufacturer’s recommendations. The blots were blocked with 3% bovine serum albumin (BSA, fraction V) in 20 mmol L⁻¹ Tris, pH 7.5, 500 mmol L⁻¹ NaCl, 0.05% Tween-20 (TTBS) solution for 3 h at room temperature. Membranes were then incubated overnight with mammalian C219 (1:1000), MRPM6 (1:50), and M2M-4 (1:50) monoclonal antibodies (Signet Laboratories, Dedham, MA, USA) in antibody buffer (1% BSA in TTBS), followed by three 5-min washes in TTBS, and a 60-min conjugate binding with a 1:3,000 solution of Goat Anti-mouse IgG horseradish peroxidase (GAM-HRP) conjugate (BioRad Laboratories, Hercules, CA, USA) in antibody buffer. Following TTBS and tris buffered saline (TBS) washes, immunoreactivity was detected using Opti-4CN Substrate Kit (BioRad, USA). Protein Ladder 11 kDa to 170 kDa (Fermentas, Canada) was used for molecular weight determination.

**Phylogenetic analysis**

All sequences were aligned using Clustal X software (24) and then adjusted by eye. The alignments were trimmed at the ends with BioEdit 7.0.5.2 to avoid inclusion of missing data (25). Phylogenetic relationships were estimated against two optimality criteria, maximum likelihood (ML) and neighbour joining (NJ) using a software program PAUP* version 4.0b10 (26). The ML starting tree was obtained via stepwise addition and replicated 100 times, with each replicate starting with a random input order of sequences. Global alignment and lookup for similar sequences were done using the BLAST Internet service (27).

<table>
<thead>
<tr>
<th>ORGANISM</th>
<th>Pgp (Accession number)</th>
<th>Mrp (Accession number)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Common carp</td>
<td>AY999964</td>
<td>FJ890347</td>
</tr>
<tr>
<td><em>(C. carpio)</em></td>
<td>198 bp</td>
<td>1425 bp</td>
</tr>
<tr>
<td>Barbel</td>
<td>DQ059069</td>
<td>FJ890350</td>
</tr>
<tr>
<td><em>(B. barbus)</em></td>
<td>197 bp</td>
<td>2091 bp</td>
</tr>
<tr>
<td>Chub</td>
<td>AY999966</td>
<td>FJ890348</td>
</tr>
<tr>
<td><em>(S. cephalus)</em></td>
<td>197 bp</td>
<td>2136 bp</td>
</tr>
<tr>
<td>Sneeep</td>
<td>AY948951</td>
<td>AY948950</td>
</tr>
<tr>
<td><em>(C. nasus)</em></td>
<td>167 bp</td>
<td>120 bp</td>
</tr>
<tr>
<td>Prussian carp</td>
<td>DQ059072</td>
<td>FJ890349</td>
</tr>
<tr>
<td><em>(C. auratus gibelio)</em></td>
<td>197 bp</td>
<td>1431 bp</td>
</tr>
</tbody>
</table>
RESULTS AND DISCUSSION

Detection of Pgp (Abcb1) and MRP (Abcc)-related sequences

Based on known full and/or partial Pgp (Abcb1) and MRPI/2 (Abcc1/2) sequences from vertebrate and invertebrate species, we designed specific pairs of primers (Table 1). Using the primer pair A/B, designed to detect the Pgp transcript, we obtained a fragment of 199 bp in length in all target cyprinids. Amplification with C/D and E/F MRP specific primer pairs resulted in over 2,000 bp fragments in all cyprinids except sneep. For sneep we obtained only a short 120 bp fragment with the G/H primer pair.

After cloning, sequencing, and computer-based sequence analysis using BLAST (NCBI), we confirmed high sequence homology at Pgp/MRP gene and protein level with other species. The obtained sequence data were submitted to the GenBank database. Identified partial Pgp mRNA sequences were registered with following accession numbers: common carp-AY999964, chub-AY999966, sneep-A948951, barbel-DQ059069, silver prussian carp-DQ059072. MRP sequence corresponded to Mrp1 (Abcc1) – barbel, Mrp2 (Abcc2) – chub and Prussian carp, and Mrp3 (Abcc3) – common carp. The related accession numbers are common carp-FJ890347, chub-FJ890348, sneep-A948950, barbel-FJ890350, and silver prussian carp-FJ890349 (Table 2).

Multiple alignments of Pgp sequences are shown in Figure 1. These sequences are homologous to Pgp genes from other organisms and share a high degree of homology with other fish and mammals (76 % to 96 %). The figure shows which nucleotides are the most conserved between different taxonomic groups.

The obtained homologies for the predicted Mrp peptide sequences among target fish species were in the range of 80 % to 95 % for Mrp1 (Abcc1; barbel), 73 % to 96 % for Mrp2 (Abcc2; chub and Prussian carp), and 80 % to 95 % for Mrp3 (Abcc3; common carp).

It was interesting to note that use of the same pair of primers resulted in fragments related to two different genes: Mrp1 (in barbel) and Mrp3 (in common carp) (Table 1). Based on these data we assume that both mRNA transcripts are present in fish liver cells and analysis of more clones could have probably resulted in the identification of both genes in each organism.
Phylogenetic analysis

The translated sequences were aligned with various ABC proteins using Clustal X software and phylogenetic trees were generated using the neighbour-joining and maximum likelihood methods. Phylogenetic analysis showed that all identified sequences grouped within the ABCB or ABCC subfamily of other organisms (Figure 2).

The analysis of family relationships through short gene and protein sequences from all tested animals clearly showed that the selected primers target highly conserved domains of P-glycoprotein. Consequently, for a more specific phylogenetic analysis, it would be desirable to obtain longer sequences.

Immunochemical detection of Pgp (Abcb1)– and MRP (Abcc)-like proteins in fish liver cells

As homologous antibodies against teleostean Pgps are not available to date, the C219 monoclonal antibody raised against human Pgp has been used in immunochemical studies to detect a conserved Pgp epitope in aquatic organisms (28, 29) within the expected range of Pgps (130 kDa to 170 kDa). Immunoblotting with C219 confirmed the presence of Pgp-related proteins in the liver of all fish species (Figure 3). Unfortunately, there are still no adequate antibodies for the detection of MRP-related proteins in fish and other aquatic organisms. Both MRPM6 (human anti-MRP1) and M2I-4 (human anti-MRP2) antibodies failed to recognise MRP-related proteins in the liver of any fish species studied (not shown), additionally supporting their already reported lack of cross-reactivity with non-human/mammalian MRPs (30).

![Figure 2](image2.png)

**Figure 2** Molecular phylogenetic relationships based on maximum likelihood (ML) analysis of our identified sequences with ABCB, ABCC, and ABCG-related sequences from other taxonomic groups. ML analysis was done by stepwise addition of WHAT and replicated 100 times, with each replicate starting with a random input order of sequences in program PAUP* 4.0b10.

![Figure 3](image3.png)

**Figure 3** Immunoblot detection of Pgp (Abcb1) in fish liver membrane vesicles (lane 2 = sneep; lane 3 = chub; lane 4 = prussian carp; 5 = barbel; 6 = common carp; 30 µg of membrane vesicle protein per lane). Membrane vesicles from male bovine cortex were used as positive control (lane 1; 10 µg of protein per lane). MM = position of molecular mass markers.
CONCLUSIONS

Multixenobiotic resistance phenotype expressed in aquatic organisms serves as a defence mechanism that protects these organisms by pumping structurally diverse xenobiotics out of the cells. In analogy to mammalian cells, MXR in aquatic organisms is mainly mediated by transmembrane ABC proteins. The best-studied ABC protein in aquatic organisms is MDR P-glycoprotein (Pgp, Abcb1), while recent studies demonstrate the protective role of other types of transport proteins such as MRPs (ABCCs) and the presence of the Mrp-related genes in fish. This study has demonstrated that (a) the MXR defence system in cyprinids, and most probably in teleost fish species in general, is mediated by Pgp (Abcb)- and MRP (Abcc)-type of efflux transporters; (b) identified sequences grouped into specific subfamilies of ABC transport proteins, revealing evolutionary closeness to Abcb1 and Abcc1-3; and (c) reliable immunochimical determination of the expression of Mrp (Abcc) proteins in fish requires development of specific antibodies. These new findings should help to better identify and characterise all proteins possibly involved in MXR in fish.

Acknowledgements

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Sažetak

ODREĐIVANJE PRISUTNOSTI EKOTOKSIKOLOŠKI VAŽNIH Pgp I MRP-PROTEINA U ŠARANKAMA NA GENSKOJ I PROTEINSKOJ RAZINI

Mehanizam multiksenobiotički otpornosti (MXR) stanični je obrambeni sustav odgovoran za svojstvo istovremene otpornosti na različite ksenobiotike, koje se očituje smanjenjem akumulacije, odnosno povećanjem izbacivanja potencijalno toksičnih tvari iz stanica vodenih organizama. MXR-mehanizam pokazuje odlike analogne fenomenu istovremene otpornosti na različite lijekove (engl. multidrug resistance, MDR) prvi put dokazanom u tumorskim stanicama. Posredovan je istim ABC transportnim proteinima kao i MDR. Istraživanja vezana uz MXR vodenih životinja uglavnom su imala težište na određivanju prisutnosti i funkcije P-glikoproteina (Pgp). Budući da se ribe iz reda šaranki često rabe u biomonitoringu slatkovodnih voda, cilj ovog istraživanja bilo je određivanje prisutnosti dvaju glavnih MDR/MXR-tipova proteina - Pgp i MRP-tip proteina - u jetri iz pet vrsta šaranki; šarana (Cyprinus carpio), klena (Squalius cephalus), mrene (Barbus barbus), bušuške (Carassius auratus gibelio) i podusta (Chondrostoma nasus). Njihova prisutnost utvrđena je na razini mRNA te na proteinskoj razini. Različiti parovi početnica dizajnirani su kako bi se identificirali homologni fragmenti gena sličnih MXR-u. Detekcija na razini proteina napravljena je putem Western blot analize s pomoću specifičnih monoklonskih protutijela proizvedenih da prepoznaju konzervirane epitope; Pgp (C219), MRP1 (MRPm6) ili MRP2 (M2I-4) u sisavaca. Transkripti obaju ključnih MXR-transportera identifičirani su u svim jedinkama, a napravljena je i filogenetska analiza dobivenih sekvenci. Imunokemijskom detekcijom s protutijelima sisavaca nismo uspjeli detektirati prisutnost MRP-proteina, dok je Pgp-ekspresija potvrđena u svih pet vrsta šaranki. Nove spoznaje da je za MXR-mehanizam zaslužno više transportnih proteina zasigurno će pridonijeti potpunijoj karakterizaciji MXR-a kao integralnog dijela detoksikacijskog, odnosno temeljnog obrambenog sustava vodenih organizama te njegovoj znanstvenoj afirmaciji kao vjerodostojnog pokazatelja kvalitete okoliša.

KLJUČNE RJEČI: ABC-proteini, filogenetska analiza, mehanizam multiksenobiotičkih otpornosti (MXR), vodeni organizmi, Western blot analiza

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