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Rainbow trout (Oncorhynchus mykiss) contain two calnexin genes which encode distinct proteins

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**Abstract**

Calnexin (IP90/P88) is an integral membrane protein of the endoplasmic reticulum that binds newly synthesized N-linked glycoproteins during their folding in the ER including MHC class I molecule. This manuscript reports the identification of two unique cDNA clones of calnexin in rainbow trout. Both encode putative mature proteins of 579 and 592 aa respectively in addition to a 24 aa signal peptide. Sequence analysis revealed that only one of the two cDNA clones encodes a putative ER retention signal, K/QEDDL, followed by a serine phosphorylation site conserved with mammalian homologs. Amino acid sequence alignment illustrated conservation of the calnexin luminal domain, which consists of a globular and a P domain, in both copies. Southern blotting revealed that there are at least two copies of the calnexin gene in the trout genome and northern blotting showed a wide tissue distribution of an estimated 3 kbp calnexin transcript with an additional minor transcript of 2.3 kbp expressed only in head kidney, spleen PBLS and strongly in RTS11. Importantly, the smaller transcript was predominantly upregulated in RTS11 after a 24 h treatment with the calcium ionophore A23187. In western blots, calnexin was detected primarily as a 120 kDa protein and upon A23187 treatment; a 100 kDa band was most prominently expressed. These results suggest that in salmonids there are two differentiated versions of the calnexin gene which encode proteins that may have diverged to perform unique biological functions.

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**1. Introduction**

Calnexin (IP90/P88) is an integral membrane protein of the endoplasmic reticulum that binds monoglycosylated oligosaccharides. It was first identified as a protein that associates with partly folded MHC class I molecules, T-cell receptors and membrane immunoglobulins (Degen and Williams, 1991; Ahiulwala et al., 1992; Hochstenbach et al., 1992). Together with calreticulin, its soluble homologue, it participates in the calnexin/calreticulin cycle responsible for folding and quality control of newly-synthesized glycoproteins before their export from the ER (Ou et al., 1993; Hammond et al., 1994; Ellgaard and Helenius, 2003). Calnexin has also been shown to play a role in ER calcium regulation (Wada et al., 1991; Roderick et al., 2000), phagocytosis (Müller-Taubenberger et al., 2001) and cell sensitivity to apoptosis (Guérin et al., 2008; Takizawa et al., 2004; Zuppini et al., 2002; Delom et al., 2007). Two distinct luminal domains of calnexin were revealed by crystallization (Schrag et al., 2001): the globular domain that contains the lectin binding site and an extended arm domain, containing two tandem proline rich motifs, which binds to ERp57 (Leach et al., 2002). The C-terminal domain is highly acidic with a protein kinase-dependent phosphorylation site that plays a possible role regulating the chaperone function (Ou et al., 1992; Chevet et al., 1999). One well investigated function of calnexin is stabilizing both major histocompatibility MHC class I and class II molecules (Anderson and Cresswell, 1994; Carreno et al., 1995). In humans, newly translated MHC class I heavy chain associates with calnexin rapidly, but it is released and replaced by calreticulin upon assembly of heavy chain with beta2 microglobulin. However, in murine cells either calnexin or calreticulin may associate with beta2 microglobulin-heavy chain dimers. Interestingly, experiments assessing the function of calnexin in the biogenesis of class I molecules showed contradicting results. Co-expression of calnexin or calreticulin with MHC class I heavy chain in Drosophila melanogaster cells demonstrated fivefold enhanced assembly with the beta2 microglobulin, while treatment with ER glucosidase I and II, inhibitors of monoglycosylated oligosaccharide formation, reduced the assembly and surface expression of MHC class I (Vassilakos et al., 1996). However, other experiments using calnexin human deficient cell lines showed no effect at all on class I assembly, transport or peptide loading (Sadasivan et al., 1995; Scott and Dawson, 1995). These experimental discrepancies may be either explained by the different model systems used or by the redundancy of molecular chaperones that can functionally replace calnexin in the ER such as calreticulin (Zhang and Williams, 2006).

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Co-immunoprecipitation studies have also demonstrated that calnexin can function as chaperone for some viral glycoproteins such as the G protein of vesicular stomatitis virus (VSV) and more recently the S glycoprotein of severe acute respiratory syndrome coronavirus by assisting in their folding and full maturation (Hammond and Helenius, 1994; Fukushima et al., 2012). Calnexin homologues have been identified in many eukaryotes such as plants, yeast, Xenopus (Huang et al., 1993; Parlati et al., 1995; Yamamoto and Nakamura, 1996) several fish species [Fuller et al., 2004, Danio rerio NM_213448.1, Takifugu rubripes XM_003978229 and XM_003978694] and remarkably in Dictyostelium (Müller-Taubenberger et al., 2001), which implies possible conserved functions for this molecule. To date, calnexin’s role in teleost fish has only been investigated in channel catfish where it was shown to associate both with the non-glycosylated MHC class II alpha chain and the glycosylated beta chain (Fuller et al., 2004). However, no studies have been carried out to assess its role during ER stress induced conditions is described.

2. Materials and methods

2.1. Fish

Rainbow trout were obtained from Silver Creek Aquaculture (Erin, ON) and kept at 13 °C in 200 L fresh-water flow-through tanks at the University of Waterloo. Animals were kept using CACC guidelines under a permit from the University of Waterloo animal care committee. Adult fish (~800 g) were anesthetized in 1 ml/L of 2-phenoxyethanol (Sigma Aldrich St. Louis, MO). Blood was drawn from the caudal sinus as previously described [Sever et al., 2013] and tissues samples were collected in RNA later (2.5 mM Na citrate, 5.3 M (NH₄)₂SO₄, 0.01 M EDTA, pH 5.2).

2.2. Rainbow trout cell lines

The rainbow trout cell lines utilized were the monocyte/macrophage cell line, RTS11 (Ganasin and Bols, 1998), and the epithelial-like cell lines: RTL-W1 from a normal liver (Lee et al., 1993), RTgill-W1 from the gill (Bols et al., 1990) and RTovarian fluid (TK Vo and Bols, unpublished). RTS11 was grown at 20 °C in L-15 media with 15% FBS and other cell lines were grown in 10% FBS as described by Kawano et al. (2010). The epithelial cell lines maintained epithelial-like morphology during the course of this study.

2.2.1. RTS11 stimulation with A23187 calcium ionophore and poly I:C

Cultures were seeded to 2 x 10⁶ in a 25 cm² flask containing L-15 media supplemented with 150 U/ml of penicillin and 150 mg/ml streptomycin in 15% fetal bovine serum (ThermoFisher Scientific, Nepean, ON). Treatment included either 2 μM of A23187 dissolved in DMSO or 50 μg/ml of polyinosinic-polycytidylic acid (poly I:C) in PBS (Sigma Aldrich, St. Louis, MO). The same volume of vehicle was added to the control.

2.3. Cloning of rainbow trout calnexin cDNA

Degenerate primers were designed based on conserved regions identified by alignment of the following sequences from GenBank: Homo sapiens [NP_001019820.1], Mus musculus [NP_031623.1], Canis familiaris [NP_001003232.1], Rattus norvegicus [NP_742005.1], D. rerio [XP_002665576.1] and Ictalurus punctatus [AAQ18011.0]. Total RNA was extracted from 2 x 10⁶ RTS11 cells using Qiagen RNeasy extraction kit according to manufacturer’s instructions (Qiagen, Mississauga, ON), followed by single strand cDNA synthesis using a Fermentas RevertAid™ First Strand cDNA Synthesis Kit with 1 μg of total RNA (ThermoFisher Scientific, Nepean, ON). PCR reactions of 25 μl included 1 μl PCR buffer, 200 μM dNTP mix, 2 mM MgCl₂, 10 μM each of the forward 5´AARTAYGAYGG NAARTGG3´ and reverse primer 5´TCTCTTTTCTAGTGCCAGCA3´, 1 μl of RTS11 cDNA and 1 μ of Taq Polymerase (MP Biomedical, Solon, OH). The reaction parameters were 95 °C for 5 min followed by 30 cycles of (95 °C 40 s, 50 °C 30 s, 72 °C 2 min) with a final 15 min 72 °C using a BioRad DNA Engine thermocycler (BioRad, Mississauga Ontario). An estimated 1200 bp PCR product was purified using a Qiagen gel extraction kit (Mississauga, ON), then ligated into pGEM-T Easy vector, subcloned into XLI-Blue MRF E. coli competent cells and sequenced at The Centre for Applied Genomics (Toronto, ON, Canada as described in Sever et al. (2013)).

2.3.1. Generation of full length cDNA

The 5’ end of the calnexin transcript was obtained using RACE adapted from Frohman et al. (1988) and described in detail in Sever et al. (2013). Briefly, cDNA synthesis used four μg of total RNA derived from peripheral blood leukocytes and gene specific primer1 (GSP1) 5’ GATGCTCCACCTCTCCA3’ and reverse transcription was performed for 60 min at 42 °C, 5 min at 45 °C, 5 min at 50 °C and 10 min at 70 °C followed by addition of RNAseH (Invitrogen, Carlsbad, CA), dATP and terminal deoxynucleotidyl transferase (Invitrogen, Carlsbad, CA) as described by Sever et al. (2013). The PCR reaction was performed using 10 μM gene specific primer sense primer 2 (GSP2) 5’ GGGGCGTGGTTGATGCTACT3’ combined with the RACE primer 5´CACTGACAGACGACTGACAGGACGCTCAAGGTTTTTTT 3’ and 2 μl of template obtained from the 5’cDNA pool. The PCR reaction for the first cycle was performed under the following conditions: 95 °C for 5 min, 48 °C for 1 min and 72 °C for 15 min with additional 30 cycles of 94 °C for 15 s, 52 °C for 30 s and 72 °C for 3 min.

Sequence alignment of the 3’UTR region of both I. punctatus NM_001200180 and D. rerio NM_213448 revealed a conserved region of 18 nucleotides which facilitated the design for a PCR primer to amplify this end of the cDNA. The primers that were used in this PCR reaction were as follows: 5’ATGGAGTTGAATGTGAGGTGTG3’ and anti-sense primer 5’AAGCTCATTGACCGACGTTTCC3’. The PCR reaction to amplify the 3’ UTR was performed under the following conditions: 95 °C for 5 min, 53 °C for 30 s and 72 °C for 4 min followed by with additional 30 cycles of 94 °C for 3 min, 53 °C for 30 s and 72 °C for 3 min.

2.4. Southern blotting analysis

Genomic DNA obtained from rainbow trout peripheral blood leukocytes was extracted using a Wizard Genomic DNA purification kit (Promega, Madison, USA). Briefly, 10 μg of genomic DNA was completely digested for 4 h at 37 °C using Fast digest enzymes EcoRV, HindIII, KpnI and PstI and BamHI (ThermoFisher Scientific, Nepean, ON), separated on a 1% agarose gel and transferred to a positively charged membrane (Roche, Manheim, Germany), followed by a UV cross linking. Membranes were washed and hybridized with DIG labelled probes as previously described (Sever et al., 2013). Two distinct probes were generated either to the proximal or distal end of calnexin. The primers used to generate the 477 bp proximal probe were as follows: forward 5’AGAGA CCTTATTTGATGATTG3’ and reverse 5’AGCCCACTTTTCAAT CCGTGGTGACCAAG3’. For the distal 540 bp probe primers were...
as follows: forward 5'TGGATGATCAGCCAGACGTACA' and reverse 5'TGCCAGAAGGAACAAAGTGAGGACG'. The bands were detected using anti-DIG antibody conjugated to alkaline phosphatase and CDP star (Roche, Mannheim, Germany).

2.5. Northern blotting analysis

RNA was isolated from both rainbow trout tissues and the RTS11 cell line using Trizol according to manufacturer’s protocol (Invitrogen, Carlsbad, CA, USA). Briefly, 8 µg of RNA from tissues and 6 µg RNA obtained from RTS11 cells were separated on a 1% agarose-formaldehyde gel and transferred to a positively charged membrane (Roche, Manheim, Germany), followed by a UV cross-linking. The membrane was washed, hybridized and detected as previously described (Sever et al., 2013). The primers used to generate the 728 bp probe were as follows: forward 5'TGGTGGTGA ACCCAGACG' and reverse 5'CACGAGCGCTGCTGCTGAC'.

2.6. Phylogenetic tree construction

Nucleotide sequences were obtained from the GenBank database (accession No.): H. sapiens CANX (NM_001024649.1), Pan troglodytes CANX (XM_003310996.2), Bos taurus CANX (NM_001105612.1), Canis lupus familiaris CANX (NM_001003232.1), Mus musculus CANX (NM_007597.3), R. norvegicus CANX (NM_172008.2), Gallus gallus CANX (NM_001030620.1), Danio rerio CANX (NM_213448.1), Caenorhabditis elegans CANX (NM_066775.3), Orzya sativa Os04g042100 (NM_001059227.1), Arabidopsis thaliana CNX1 (NM_125573.3), Xenopus laevis CANX-a (NM_001086857), Xenopus tropicalis CANX (NM_001005668), D. melanogaster CNX99A (NM_170407.3), T. rubripes LOC10107085 (XM_003978229), T. rubripes LOC101069714 (XM_003978694), Orzyas latipes LOC101156199 (XM_00409293), O. latipes LOC101159642 (XM_004056545), I. punctatus LOC100304606 (NM_001200180.1) and Oreochoemis niloticus LOC100107166 (XM_003447644). The alignment was generated using Muscle (Edgar, 2004) and was used to construct a phylogenetic tree using the neighbour-joining method (Saitou and Nei, 1987) with the Jukes and Cantor correction (Jukes and Cantor, 1969) for the per cent bootstrap confidence values.

2.7. Sequence analysis

SignalP 3.0 server was used to identify the signal peptide (Bendtsen et al., 2004), while TMHMM Server v. 2.0 was used to identify transmembrane regions (Krogh et al., 2001), NetPhos 2.0 phosphatase (Sigma Aldrich St. Louis, MO). Bands were detected using NBT/BCIP (Roche, Manheim, Germany) according to manufacturer’s instructions.

2.8.1. Isolation of peripheral blood leukocytes

Trout peripheral blood was centrifuged at 400g for 10 min, a buffy coat was obtained and diluted 1:10 with L-15 media (Sigma Aldrich St. Louis, MO). Diluted Fractions were layered on the top of 3 ml histopaque 1077 (Sigma–Aldrich, St. Louis, MO) and centrifuged for 30 min at 400g. White blood cells were collected, centrifuged and washed once with 1× phosphate buffered saline (PBS) for 5 min. Pellets were stored in −80 °C until later use.

2.9. Immunofluorescence

RTS11 cells were seeded to 2 × 105 cells per well in 4 chamber slides containing 2% PBS 150 U/ml of penicillin and 150 mg/ml streptomycin and grown overnight. All steps were conducted at 20 °C. Cells were washed once with PBS (137 mM NaCl, 2.7 mM KCl, 4.3 mM Na2HPO4, 7H2O, 1.4 mM KH2PO4; pH 7.4) and fixed for 15 min with 4% paraformaldehyde (Thermo Fisher Scientific, Nepean, ON) followed by two washes with PBS. Cells were permeabilized by incubation for 10 min with PBS containing 0.25% Triton X-100 (PBST), followed by 3 washes with PBS of 5 min each. Cells were incubated with 1% BSA in PBST containing 0.1% (v/v) of Triton X-100 for 30 min and then probed with 1:100 of polyclonal rabbit anti-human calnexin (GTKX101676) in 1% BSA for 1 h. After three washes for 5 min each in PBS, cells were incubated for 1 h with 1:3000 of goat anti-rabbit Alexa Fluor 488 (Invitrogen, Carlsbad, CA) in 1% BSA followed by 3 washes with PBST of 5 min each. Cells were dried for 5 min in room temperature, mounted with Fluoroshield containing DAPI (Sigma Aldrich St. Louis, MO), coverslips were attached and the slides were stored at 4 °C. Slides were examined by laser scanning confocal microscopy using a Zeiss Axiovert 200 microscope and ZEN 2009 software (Carl Zeiss, Mississauga, ON).

3. Results

3.1. Sequence analysis

An estimated 1230 bp amplicon was isolated from RTS11 cDNA which showed 83% nucleotide identity with I. punctatus calnexin (NM_001200180.1). In order to obtain the 5’end and 3’end of the trout calnexin cDNA, PCR primers were used for 5’ RACE and for a conserved region in the 3’UTR using a PBL cDNA template. The corresponding PCR fragments were 380 bp and 320 bp respectively. PCR amplification of the full open reading frame of rainbow trout calnexin was then conducted using 5’ and 3’ gene specific primers using PBL cDNA template. Interestingly, two unique cDNA clones similar in length were obtained.

The first cDNA clone contains 1812 nt in its coding region and 94 nt of 3’UTR corresponding to putative 603 aa protein of 68 kDa. The second cDNA clone contains slightly longer sequence of 1851 bp with additional 23 bp of 3’UTR (117 bp total) which encodes a 616 aa protein with predicted molecular weight of 70 kDa.

Both cDNAs showed 73% and 79% nucleotide identity with H. sapiens and I. punctatus Calnexin sequences, respectively with 72% and 75% sequence identity to their encoded proteins. The main characteristic of the rainbow trout calnexin proteins are illustrated in Fig. 1. The N terminal domain includes a 24 aa signal peptide, followed by a luminal domain which contains a globular domain and an extended arm domain with a central segment, the P domain. This segment contains two proline motifs: M1 and M2 repeated in tandem 4 times each with the consensus sequences of M1: I-DP(D/E)A-KPEDWD(D/E) and M2: G-W-P-IN-P-Y. Moreover,
there are 3 regions that have high sequence identity with human calnexin that are named box a, b and c. The box a segment is located upstream to calreticulin domain 1 whereas box b and c flank the repeated motifs. There is also a transmembrane domain of 21 aa that was predicted by TMHMM Server. Interestingly, there are two pairs of cysteines predicted to be palmitoylated by CSS-Palm 3.0 immediately after the transmembrane domain. These cysteines are found to be conserved across many species and when palmitoylated were shown to be critical for the association of calnexin with the ribosome translocon complex, allowing calnexin to bind to the N-linked glycans as they exit from the translocon (Lakkaraju et al., 2012). Another shared feature with mammalian calnexin is the presence of an acidic cytoplasmic tail enriched with glutamic acid and ER retention signal KQEDDL, which appears only in one of the clones and interestingly does not appear in the channel catfish sequence (Fuller et al., 2004). There are 5 phosphorylation sites predicted by NetPhos 2.0 Server for both trout clones, however a unique serine phosphorylation site downstream of the ER retention signal known to regulate calnexin activation (Chevet et al., 2010) appears only in calnexin cDNA sequence containing the ER retention motif.

3.2. Phylogenetic tree

In order to further examine the evolutionary relationship between the unique trout clones and other calnexin nucleotide sequences, a phylogenetic tree were constructed (Fig. 2). As expected, strong distinct clusters can be seen for the tetrapod, teleost fish and plant sequences. Both trout sequences clustered together with a short genetic distance suggesting a recent gene duplication event, which is common in salmonids (Schmidtke and Kandt, 1981). Interestingly a more ancient duplication event also seems to have occurred in teleost fish, as both pufferfish, and rice fish showed a second sequence which grouped with zebrafish calmegin outside the main cluster of fish, frog and mammalian calnexin sequences, but still within a vertebrate cluster. This suggests a very early duplication event to produce this testis-specific ER chaperone homologous to calnexin.

3.3. Trout calnexin gene copy number

To further examine the possibility that the calnexin gene might be duplicated in trout, two separate probes were designed to proximal and distal segments of the calnexin gene. When the proximal probe was used, between 3 to 4 bands were observed in each digest (Fig 3A). The distal probe showed similar results showing two dark bands and either one to two faint bands (Fig 3B). These results suggest that the calnexin gene is present in at least two copies in trout, unlike its soluble paralog calreticulin which appears in rainbow trout as single copy gene (Kales et al., 2004).

3.4. Calnexin tissue expression

Northern blotting shows wide tissue distribution of the calnexin mRNA in trout tissues with the most prominent message being an estimated 3 kb transcript band (Fig. 4A), but with a faint lower band detected in spleen, head kidney and PBLs. This lower band is strongly expressed in RTS11 (Fig 4B). The highest expression of that transcript was observed in spleen, head kidney PBLs (Fig 4A) and in the cell line RTS11 (Fig 4B). In order to determine if both bands were similarly regulated, RTS11 cells were treated with A23187, which is known to induce ER stress and upregulate other molecular chaperones in vitro such as ERP57 (Sever et al., 2013). Importantly, the lower calnexin transcript was upregulated up to 2.6-fold in 24 h followed by a decreased expression at 36 h. Interestingly, the upper band expression remained relatively constant (Fig 4B and C). This result points out to the possibility that trout may have two unique transcripts responding differentially to ER induced stress due to unique modes of regulation.

3.5. Calnexin protein expression under normal and induced ER stress conditions

In order to further explore the possible role of calnexin during ER stress, an anti-human calnexin antibody (GTIX101676) was used. This antibody was generated against the recombinant P domain of human calnexin, which shares 81% identity with the trout calnexin sequence. As opposed to its 70 KDa predicted molecular weight, trout calnexin protein was detected as a 120 kDa band in RTS11, RTgill, RTBUt, RTovarian fluid and RTliver (Fig. 5A and B). Interestingly, a lower fainter band around 100 KDa was also detected in all cell lines tested. A doublet band around 55 kDa was observed in RT liver, probably due to antibody binding to calreticulin, known to migrate as a doublet band in trout (Kales et al., 2007). In channel catfish calnexin is detected as a single band around 100 KDa compared to its 69 kDa molecular weight (Fuller et al., 2004). This discrepancy is common in many species and might be explained by the abundance of acidic amino acids which interferes with SDS binding and reduce the mobility on SDS–PAGE, and post-translational modifications. Importantly, induction of RTS11 with A23187 in a set of three repeats showed detection of the lower band in the treated cells but not in the control (Fig 6). RTS11 treated with poly IC did not show any induction of the lower band (data not shown) which suggests that the expression of the smaller version of calnexin is modified specifically under ER stress conditions.

3.6. Calnexin localisation in RTS11 cells

In order to further validate the specificity of the GTIX101676 antibody, RTS11 were stained with anti-calnexin antibody (GTIX101676) and examined under the confocal microscope (Fig. 7). The staining revealed a meshwork structure within the cells, with additional staining observed surrounding the nucleus. A similar and stronger staining pattern was observed for other ER resident proteins, trout ERp57 and calreticulin the latter of which contains a putative ER retention signal (Kales et al., 2004). These findings further support the localisation of calnexin to the ER membrane.

4. Discussion

In this study, rainbow trout calnexin was characterised and its expression was investigated to examine the regulation of a possible peptide loading complex in fish using rainbow as a model organism. Two unique cDNA clones encoding trout calnexin which share 99% sequence identity in their coding region were identified. Intriguingly, only one cDNA version contains a unique c-terminal with putative ER retention signal and conserved serine phosphorylation site which activates calnexin when protein misfolding occurs and is important for the recruitment of calnexin to ribosome translocons (Cameron et al., 2009; Chevet et al., 1999; Lakkaraju et al., 2012). Sequence alignment with the human calnexin protein reveals 72% identity and implies structural and functional conservation of calnexin as an ER lectin chaperone, calcium binding protein and as a protein indirectly assisting in oxidative folding. The luminal domain of calnexin contains the two conserved domains: the globular and the arm domain. The lectin binding site was found in a pocket on the surface of the globular domain (Schrag et al., 2001) whereas the tip of the arm domain contains the docking site that recruits ERP57 (Leach et al., 2002) and therefore helps to
Fig. 1. An alignment of rainbow trout calnexin protein sequences with other known vertebrate sequences obtained from the GenBank (see Section 2 for specific accession numbers). Numbers on the right indicate amino acid positions. The conserved domains are labelled in bold above the sequences. The palmitoylation site in the juxta-membranal region is underlined. Both the putative ER retention signals and the serine phosphorylation site are labelled in bold and underlined.
support oxidative folding in the ER. The P domain in the central arm segment contains two conserved motifs repeated four times each which contain low capacity and high affinity calcium binding sites, while the c-terminal domain is highly enriched with acidic amino acids shown to be as high capacity low affinity calcium binding sites (Michalak et al., 1992). All of these features are evident in the trout calnexin sequences, suggesting that many of its roles are conserved in trout, however the differences in the C terminal domains between the calnexin cDNA sequences might suggest of unique mode of regulation and perhaps differential subcellular localisation.

A phylogenetic tree constructed with calnexin nucleotide coding sequences indicates that the two copies arose in a recent gene duplication event within trout, or more probably salmonids (Schmidtke and Kandt, 1981) similar to that seen for trout ERp57 (Sever et al., 2013), suggesting that both are bone fide calnexin homologues. A duplication of calnexin like sequences of pufferfish and rice fish obtained from different loci may be more ancient as these sequences cluster outside all of the vertebrate calnexin sequences. Sequence alignment of their corresponding proteins reveals sequence identity of 73% and 84% respectively to zebrafish calmegin, a calnexin homologue in the testis which is expressed exclusively during spermatogenesis both in mouse and human calnexin and was suggested to bind sperm surface proteins important for egg-sperm interaction (Watanabe et al., 1994; Tanaka et al., 1997; Ikawa et al., 1997).

In order to investigate the regulation of calnexin during ER stress, cells were treated with calcium ionophore A23187 which has been previously shown to upregulate calnexin transcript expression by up to 2.5-fold in S. pombe and appeared as a single band in a northern blot (Parlati et al., 1995). Interestingly, in trout two possible transcript bands for calnexin were observed that are expressed only in immune system organs. Surprisingly, ER stress induction in RTS11 using A23187 showed that the smaller, less abundant transcript band was induced in contrast to the main transcript which remained unchanged. Similar results under the same stimulating conditions were obtained at the protein level showing the induction of a less abundant 100 kDa band with no change in expression of the higher molecular weight protein band. These results support the presence of two trout genes for calnexin that are regulated differently under ER stress conditions and perhaps performing unique biological functions. Importantly, according to their predicted cDNA sequences their difference in size might be more due to post translational modification rather than simply amino acid differences.

In contrast to what was seen here with trout calnexin, trout calreticulin, the soluble homologue of calnexin known to perform similar functions in the ER, was identified as a single copy gene in trout and failed to upregulate under several ER stress conditions (Kales et al., 2007). This suggests that fish calnexin may perform functions during ER stress that calreticulin cannot. Under normal conditions, trout calreticulin expression was found to be highest in the liver (Kales et al., 2007) while trout calnexin seems to be more abundant in immune system organs. Interestingly, while calnexin deficient mice display impaired motor function they are not embryonic lethal (Denzel et al., 2002) while calreticulin deficient mice exhibit an embryonic lethal phenotype due to a defect in heart development (Michalak et al., 2002). In addition, it has been recently demonstrated that both genes are important for generation of neuromasts during lateral line development in zebrafish using calnexin and calreticulin knockout morpholinos, however the calnexin knockout had a more profound effect on this develop-
ment (Hung et al., 2013). These results suggest that both in mammals and in fish calreticulin and calnexin play a role during embryonic development, which suggest these two proteins may have distinctive roles outside of their chaperone function. Monitoring the expression of these two calnexin genes during stages of embryogenesis in rainbow trout could shed a light on their possible unique functions during development.

The immunocytochemistry of calnexin in RTS11 suggests that calnexin is localised to the ER and possibly resides close to the nucleus, as in some cells it showed staining near the nucleus. Interestingly, calnexin has been found as the major calcium binding protein in rat hepatic nuclear membranes although appeared as a minor member of the total nuclear protein (Gilchrist and Pierce, 1993). In addition, immunostaining of calnexin in rat epithelial cells has demonstrated a reticular network structure and nuclear envelope staining (Wada et al., 1991) supporting our findings. It may be that the trout calnexin without the ER retention signal can escape the ER and perhaps be localised in the nuclear periphery where it can regulate calcium levels in this region. However, further experimentation with monoclonal antibodies to each specific form is required to see which form(s) is localised to the nuclear region.

The role of calnexin as a chaperone has been widely investigated in mammals, however some research conducted in yeast and the microorganism Dictyostelium shed a light on the other important cellular functions of calnexin such as during apoptosis and phagocytosis. The first report linking calnexin to apoptosis

Fig. 3. Southern blots of the rainbow trout calnexin gene. Left panel, labelled (A) shows the results obtained with a probe spanning the proximal domain of calnexin, right panel (B) using a probe for the distal coding domain of calnexin. The position of the size markers are shown by arrows on the right margin.

Fig. 4. Northern blots of rainbow trout calnexin showing tissue distribution of calnexin mRNA in normal rainbow trout tissues (4A) and in RTS11 upon ER stress stimulation using 2 μM Calcium ionophore A23187 for 36 h (4B). Densitometry analysis of the RTS11 experiment was conducted using image J software (4C). The estimated size of the 28s and 18s ribosomal bands is shown by the arrows on the right margin.

Fig. 5. Distribution of calnexin protein in trout cell lines using an anti-human calnexin antibody. Lysates were separated on a 10% (Fig. 6A) or 12% (Fig. 6B) SDS PAGE gels. The position of the size marker is shown on the right margin.

Fig. 6. Upregulation of Calnexin following in vitro stimulation of RTS11 with Calcium Ionophore A23187 for 24 h. This experiment is representative of the results in a three replicate set.
was conducted in an S. pombe yeast two hybrid system demonstrating that calnexin is required for cell death mediated by the interaction of its cytosolic tail with the a pro-apoptotic protein BAK (Torgler et al., 1997). Later studies in mammals using calnexin-deficient cells further demonstrated its possible role in mediating apoptosis as these cells showed resistance to ER stress induced apoptosis (Zuppini et al., 2002; Groenendyk et al., 2006). Another surprising function for calnexin came from a study with Dictyostelium, the only microorganism containing both calnexin and calreticulin proteins. This study showed that co-disruption of calnexin and ERp57 proteins attenuates the release of partially misfolded alpha1-antitrypsin to the secretory pathway. J. Biol. Chem. 284, 34350–34359.

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