Analysis of Htra Gene from Zebrafish (*Danio Rerio*)

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Abstract

HtrA which is characterized by the combination of a trypsin-like catalytic domain with at least one C-terminal PDZ domain is a highly conserved family of serine proteases found in a wide range of organisms. However the identified HtrA family numbers varies among species, for example the number of mammalian, Eschericia coli, fruit fly-HtrA family are 4, 3 and 1 gene respectively. One gene is predicted exist in zebrafish. Since no complete information available on zebrafish HtrA, in this paper zebrafish HtrA (zHtrA) gene was analyzed. The zHtrA is belonged to HtrA1 member and predicted encodes 478 amino acids with a signal peptide, a IGF binding domain, a Kazal-type inhibitor domain in the up stream of HtrA-bacterial homolog. At the amino acid sequence the zHtrA showed the 69%, 69%, 68%, 54% and 54% with the rat HtrA1, mouse HtrA1, human HtrA1, human HtrA3 and mouse HtrA4 respectively. The zHtrA1 is firstly expressed at 60 hpf and mainly in the vertebral rudiments in the tail region.

Introduction

HtrA which is characterized by the combination of a trypsin-like catalytic domain with at least one C-terminal PDZ domain is a highly conserved family of serine proteases found in a wide range of species including microbes, plants, and animals (Pallen and Wren, 1997; Clausen et al., 2002). The over 180 members of this family indicates that all sequences have a length around 350-450 amino acid residues. The number of HtrA family members were varies among species. Four, three and two HtrA members were identified in human; Eschericia coli and Synechocys; Arabidopsis thaliana respectively. Only one HtrA identified in Saccharomycases cerevisae, Schizosaccharomycoses pombe, Candida albicans and Drosophila melanogaster. In zebrafish (*Danio rerio*) one HtrA was predicted (Clausen et al., 2002).

From mammalian at least 4 members of HtrA, from HtrA1 to HtrA4 have been identified. The HtrA2 has mitochondrial signal and a unique transmembrane at N-terminal of bacterial HtrA-like. The human htrA2 is expressed in placenta and pancreas of normal human tissues and highly expressed in promyelocytic leukemia HL-60, chronic myelogenous leukemia K-562, Burkitt lymphoma Raji and human colorectal carcinoma SW480 cell lines (Facio et al., 2000) and Omi is upregulated in mammalian cells in response to stress induced by both heat shock and tunicamycin treatment (Gray et al., 2002).

Mammalian HtrA3 have characteristic structural motifs which are shared with HtrA1. In contrast to HtrA2, those HtrAs are secretory proteins having a signal sequence for secretion at the N-terminus. Those HtrAs also contain insulin-like growth
factor binding domain (IGFB), a Kazal type serine protease inhibitor domain at N-terminal regions of bacterial-HtrA homology region. The expression of HtrA3 or pregnancy-related serine protease (PRSP) gene was low before pregnancy, but it was increased at implantation and markedly up-regulated post-implantation. In-situ hybridization localized low levels of mRNA expression to the epithelium and stroma during very early pregnancy, but high expression to the decidual cells on day 8.5, primarily at the mesometrial pole where the placenta was forming (Nie et al., 2003). The HtrA1 (L56 or PRSS11) was originally isolated in differential screening for mRNAs repressed by SV40 transformation of human fibroblasts (Zumbrun and Trueb, 1996). HtrA1 was also independently isolated as a gene up-regulated in cartilage of human osteoarthritis patients (Hu et al., 1998) and aged joint cartilage (Ly et al., 2002). The HtrA1 mRNA was also known to be down-regulated in ovarian cancer (Shridhar et al., 2002), malignant melanoma, (Baldi et al., 2002), primary brain tumors, and several cancer cell lines (Chien et al., 2004). This tumor suppressive activity of HtrA1 was largely dependent on the protease activity (Hu et al., 1998). Interaction of HtrA1 PDZ domain with partner protein regulates the proteolytic activity (Murwantoko et al., 2004).

HtrA is found in many different species. Four and three member of HtrA have been reported in human or mammalian and *E. coli* respectively, but only one member has been identified in fruit fly and one member was predicted in zebrafish (Clausen et al., 2002). In this paper we identified the zebrafish HtrA member and determined the expression.

**Materials and Methods**

**Primer and amplification**

The searching of amino acid homology of mouse HtrA1 to zebrafish data base indicated that two short fragment of cDNA showed high homology. The fj28a11 showed 65% identities to IGFB and KI domain. The fk91g02 showed 81% identities with serine protease and PDZ domains.

![Amino Acid Homology](image)

**Figure 1.** The homology between two fragments of zebrafish cDNA with mouse HtrA1 was presented. The relative position of Zeb-S8-2F (2F), Zeb-S8-2R (2R), Zeb-S8-3F (3F), Zeb-S8-4F (4F), Zeb-S8-3R (3R) and a set of 5’RACE (5’Rc) primers were shown. S=signal peptide, IB=IGF binding protein, KI=Kazal-type protease inhibitor, PDZ=PDZ domain

The primers were designed to amplify zebrafish Fj fragment, region between Fk and Fj fragments, 5’ terminal end. The pair primer Zeb-HtrA-2F (ATGGATCCGAGTTGAGAAGGTCTGGAA) and Zeb-HtrA-2R (CTCTTAAGGGC TACCACATGACCATAG) were used amplify IGFBP-KI domains homolog. To amplify region between Fk and Fj region the sense primers ZEB-S8-3F (TATCGAATTCCTCATGTGGTAGCCAACA) and ZEB-S8-4F (GTGGAGTTCTGCTCTGGTTTTGTGGTCT) were designed from the 3’-end of fj fragment and one anti sense primer ZEB-S8-3R (TCTACT ATAGGTCTGATGGAATGGCAAG) was designed from 5’-end of fk fragment.

To identify the 5’ part of expected Zebrafish HtrA gene, the rapid amplification of 5’ cDNA Ends (5’ RACE) was done using Takara 5’-RACE Core Set (Takara Biomedicals). The 5’ phosphorylated RT primers (CTGAAGCTTCTCGT) was used to
synthesis first strand cDNA at 50°C for 1 h. Degradation of hybrid RNA by RNAsEH was done at 30°C for 1 h followed by circularization/ concomitamerization at 16°C for overnight. The two primer pairs (Sense 1 GTGTGCCTGTGCAAGAGGTTC Antisense 1 CGGTAGTCGTTGCACCAAC C; Sense 2 GTGTCCTACAGGAACATCTG Antisense 2 CATTCCAGACCTTCTCCACTC) were used to amplify the longer and nested unidentified sequences respectively. The nested DNA fragment was sequenced. The determination of the 3’ end zHtrA was done by analysis on genomic sequences.

RNA isolation and cDNA synthesis
RNA was isolated from different stages of zebrafish using ISOGEN (Wako Pure Chemical Industries, Osaka, Japan) following manufacturer’s instructions. In brief, the embryos were homogenized in Isogen followed by incubation at room temperature for 5 min. The solutions were extracted with chloroform and incubated at RT for 3 min. The supernatant fraction were precipitated with isopropanol and washed with 70% ethanol. Precipitated RNA was dissolved in DEPC-treated TE. One μg of total RNA was used to synthesize first strand cDNA using random primers at 37°C for 60 min (Pharmacia cDNA synthesis kit).

HtrA expression by RT-PCR
To identify the stage of fish expressing the HtrA, the 30 hpf, 60 hpf and adult stages of zebrafish cDNA were used for PCR. The pair primer Zeb-HtrA-2F and Zeb-HtrA-2R which predicted to amplify IGFBP-KI domains homolog were used. The amplification was performed 95°C (30 s), 55°C (30 s), 72°C (1 min) for 30 cycles. This fragment also cloned into pBSKSII (Stratagene).

Whole mount in situ hybridization
Recombinant plasmid of PBSKSII containing DNA fragment amplified using zeb-HtrA-2F/ zeb-HtrA-2R primers was used in this experiment. Sense and antisense digoxigenin-labeled RNA probes were synthesized with a kit (Roche Diagnostics Heideberg, Germany) using T3 and T7 RNA polymerases (Boehringer Mannheim).

The various stages of zebrafish were fixed in 4% paraformaldehyde in PBS at 4°C for overnight. After removing the vitelline membrane, the fixed embryos were dehydrated in methanol and stored at -20°C. The embryos were rehydrated in PBS and the 70 hpf embryos were treated with 50 μg/ml proteinase K for 5 min. The embryos were further fixed in 4% paraformaldehyde in PBS for 20 min. After two washes with PBST for 5 min each, the embryos were incubated in a prehybridization solution (50% formamide, 5xSSC and 0,1% Tween, 500 μg/ml yeast tRNA, 50 μg/ml heparin) at 56°C for 1 h. Hybridization was carried out at 56°C overnight in the prehybridization solution supplemented with 3 μg/ml digoxigenine-labelled probe. After hybridization, excess probes were removed by washing in 2xSSC/ 50%formamide at 56°C for 1 h and by washing in 2XSSC at RT for 10 min twice. The embryos were washed in RNase buffer (0.5 M NaCl, 10mM Tris-Hcl (pH 8.0) contain 0.1% Twen 20 then treated with 20 mg/ml RNase  in RNase buffer at RT for 10 min. The embryos were washed with 2xSSC at RT and with 2XSSC/50% formamide, 2XSSC and 0.2xSSC at 55°C for 1 h, 15 min and 15 min respectively. Finally the embryos were washed with PBST at RT for 5 min. The embryos were then treated with 1% blocking reagent (Boheringer Mannheim), containing 0.2% Twen 20 for 1 h and incubated overnight at 4°C with 1:4000 diluted of alkaline phosphatase-conjugated antidigoxigenin antibody in the blocking reagent. After extensive washing, the color
was developed using NBT/BCIP. The embryos were fix in 4%PFA, 0.25% glutaraldehyde in PBS and destaining using acetone.

Genomic analysis
Genomic analysis using Genetyx program was used to predict amino acid sequences encoded by zebrafish HtrA. Comparison the sequences with the data in gene bank was done usin BLAST in NCBI. Prediction protein structure was done in NCBI protein structure. The comparison of the predicted amino acid sequences of zebrafish HtrA which other member was done using Clustal W (Thompson et al., 1994)

Result
Identification of zebrafish HtrA
The amplification using zeb-HtrA-2F/ zeb-HtrA-2R primers using adult cDNA as template gave result a fragment 430 bp in size. This size was as been predicted since the primer will amplify from nucleotide 370 to 800. This result showed that the cDNA contains Fj fragment. To verify the present of Fk fragments and whether those Fj and Fk fragments came from one gene, the two sense primers ZEB-S8-3F, ZEB-S8-4F which designed from the 3'-end of fj fragment and one anti sense primer ZEB-S8-3R which designed from 5'-end of fk fragment were used to amplify the cDNA in separate tube. The amplification produced single fragment on every primer pair. The DNA fragment which is produced by ZEB-S8-3F/ ZEB-S8-3R primer was larger than fragment of ZEB- S8-4F/ ZEB-S8-3R primer, and at the fragments size were as been expected. Those results indicated that the Fk fragment was present in cDNA, and the Fj and Fk fragments were came from one gene. Next we identified entirely HtrA gene by RT PCR for the middle part, and by 5'-RACE for 5' terminal ends. The comparison between Fk fragment and zebrafish genomic data bank resulted the 72 nucleotide in 3' terminal ends. The sequence of entirely zebrafish HtrA gene and predicted amino acid sequences were presented in Figure 2.

Zebrafish HtrA sequences contained a clear 1330 nucleotidet (nt) ORF, with the start codon ATG at nt 139±141 and stop codon TGA at nt 1467+1469; outside the ORF is a 5' UTR 138 nt. The identical N-terminal ends of protein contain a predicted signal peptide (aa 1±22). A search of the NCBI protein structure database showed that the long-form protein contains aa sequences consistent with four other domains (Figure 1) : (i) an insulin-like growth factor (IGF)-binding domain (aa 27±91), (ii) a Kazal-type Serine protease-inhibitor domain (aa 109±151), (iii) a trypsin protease domain (aa 167±360) and (iv) a PDZ domain (aa 379±476).

A comparison of the sequence (Figure 1) with entries in the GenBank databases (April 2004) showed the greatest homology were the mammalian HtrA1 proteins. At the amino acid sequence, the zHtrA1 has 69% identity of the sequence with the rat and mouse HtrA1 (accession numbers NP_113909, AAD52683, AD52682 and AAH13516), 68% identity with human HtrA1 (NP_002766), 54% with human HtrA3 (NP_444272) and 54% with mouse HtrA4 (XP_284398). The homology alignment of those sequences is presented in Figure 3.
Figure 2. The mRNA zebrafish HtrA sequences and their deduced amino acid sequences. The start codon (ATG) are in bold. The active serine protease GNSGGPL and the additional TNAHV histidine residues are underlined and in bold. The catalytic triad amino acid His, Asp and Ser are circled. The lower case sequences was predicted from genomic sequence.
Figure 3. Amino acid sequences alignment of zHtrA, mHtrA1, hHtrA3 and hHtrA4.
The HtrA1 proteins are homologous with the HtrA from bacteria. These HtrA proteins belong to a family of serine proteases that possess the aa sequence motif of GNSGGAL (in bacteria) or GNSGGPL (in mammals) in their active sites. In addition, they display a motif of TNAHV residues in the vicinity of GNSGGPL and the triad of histidine, aspartic acid and serine for their catalytic activity (Zumburn and Trueb, 1996; Hu et al., 1998). All of these motifs are present in protein (Figure 2, 3). The serine protease active site sequence GNSGGPL is at aa 321±328, the TNAHV residues are at aa 213±216, and the catalytic triad histidine, aspartic acid and serine are at aa 216, 246 and 3245 respectively. This confirmed that the zebrafish HtrA represents an additional member of the HtrA serine protease family as Zebrafish HtrA1. Clausen et al. (2002) noted that one member of HtrA gene family is predicted in zebrafish, HtrA2, because lack of signal peptide. Here we reported that zebrafish also has a member of HtrA family which has signal peptide, i.e. zHtrA1.

Expression of zHtrA1

To observe the expression of zHtrA1, the RT-PCR and whole mount in situ hybridization was done. ZHtrA1 is not expressed at early stage of embryo as result from in situ that no signal was detected in 8 cells, 50% epiboly, 90% epiboly, 14 hpf and 25 hpf which also supported by RT-PCR result that at 30 hpf did not amplify DNA fragment (Figure 4, 5) The HtrA1 seems to be expressed at least at 60 hpf, as indicated by positive result from RT-PCR and appearance of signal on in situ hybridization at this stage. Whole mount in situ hybridization showed that Zeb-HtrA1 was detected clearly only in the vertebral rudiments in the tail region (Figure 5).

Discussion

Organization of HtrA

HtrA is a highly conserved family of serine proteases found in a wide range of
species including microbes, plants, and animals. The number of those gene’s members were varies among species. The number seems not correlated with the complexity of organisms. Since Clausen et al. (2002) notes that unicellular organism, E. coli at least has 3 members; the mammalian has at least 4 members. The fruit fly, D. melanogaster and S pombe which their complexity higher than bacteria and lower than mammalian, only have one HtrA1 member. Our screening from D. melanophila cDNA library supported that only one gene found in this animal (data not shown).

Since the HtrA is distribute wide range of organisms, thus this gene is predicted exist in zebrafish. In this study we identified zebrafish HtrA member gene, zHtrA1. However this gene is different with the predicted by Clausen et al. (2002) which belong to HtrA2 member due to the lack of signal sequence in N-terminal. It still does not know about the actual number(s) of HtrA gene in zebrafish. To address this question, the extensive screening should be carried out.

HtrA expression

HtrA1 has similar structure with HtrA3, HtrA4. Structure of N-terminal regions of mammalian HtrA1 and 3 are contain secretory signals at the N-terminus followed by two domains; one similar to the insulin-like growth factor (IGF) binding protein domain and the other to the Kazal-type serine protease inhibitor domain (Clausen et al. 2002). Although this N-terminal part did not show cross immunoreaction between HtrA1 and HtrA3 (data not shown), those two genes seem have similarity in expression. HtrA1-3 are firstly expressed in later stage of embryo. The mHtrA1 and mHtrA3 were firstly expressed at 10.5 dpc (Oka et al 2004) and at 9.5 dpc (Tocharus et al., 2004) respectively. The zHtrA1 seems to be expressed at least at 60 hpf, as indicated by positive result from RT-PCR and appearance of signal on in situ hybridization at this stage (Figure 4, 5).

The HtrA1-3 seem to be expressed in the tissues closely associated with skeletal system, as showed in mouse HtrA1 (Oka et al., 2004), mouse HtrA3 (Tocharus et al. 2004). Our result support that note in different animal as zHtrA1 was detected clearly only in the vertebral rudiments in the tail region as judged by whole mount in situ hybridization (Figure 5). The expression in the skeletal system is closely related with TGF-β expression (Oka et al., 2004), then the HtrA1 and HtrA3 have been provided their function as inhibitor signaling TGF-β family protein (Oka et al., 2004; Tocharus et al., 2004). However, in zebrafish the HtrA1 and TGF-β are firstly expressed at is different in stage. The TGF-β family member in zebrafish have function in posterior and ventral development and are firstly expressed at embryonic shield or gastrulation (Hwang et al., 1997; Lele et al., 2001; Dickmeis et al., 2001). On the other hand zHtra1 is firstly expressed at 60 hpf (Figure 4, 5). The late expression of HtrA compare the BMP seems also occurred in the mouse. At first the both HtrA3 and -1 were not expressed in the core of the cartilaginous condensations. Interestingly, however, when the blood vessels invaded into the condensations and ossification started in the peri- and postnatal periods, the expression of HtrA3 and -1 were tremendously upregulated. Chondrocytes probably undergoing degeneration in the ossification center produced HtrA1 and -3 (Oka et al., 2004; Tocharus et al., 2004).

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