Molecular cloning and expression analysis of a zebrafish novel zinc finger protein gene \textit{rfn141}

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Abstract

\textit{ZNF230} is a novel zinc finger gene cloned by our laboratory. In order to understand the potential functions of this gene in vertebrate development, we cloned the zebrafish orthologue of human \textit{ZNF230}, named \textit{rfn141}. The cDNA fragment of \textit{rfn141} was obtained by rapid amplification of cDNA ends (RACE). The open reading frame (ORF) encodes a polypeptide of 222 amino acids which shares 75.65\% identity with the human \textit{ZNF230}. RT-PCR analysis in zebrafish embryo and adult tissues revealed that \textit{rfn141} transcripts are maternally derived and that \textit{rfn141} mRNA has a broad distribution. Zygotic \textit{rfn141} message is strongly localized in the central nervous system, as shown by whole-mount \textit{in situ} hybridization. Knockdown and over expression of \textit{rfn141} can induce abnormal phenotypes, including abnormal development of brain, as well as yolk sac and axis extension. Marker gene analysis showed that \textit{rfn141} may play a role in normal dorsoventral patterning of zebrafish embryos, suggesting that \textit{rfn141} may have a broad function during early development of vertebrates.

Key words: \textit{rfn141}, zebrafish (\textit{Danio rerio}), development, zinc finger protein.

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Introduction

The zinc finger gene family, one of the largest gene families in mammals, is defined by a conserved cysteine and histidine rich domain essential for the binding of zinc ions (Freemont 1993; Klug and Schwabe, 1995). This gene family can be divided into several subfamilies, including ring finger, C2H2, glucocorticoid receptor, GATA1, GAL4, and LIM (Barlow \textit{et al.}, 1994; Borden and Freemont, 1996; Hammarstrom \textit{et al.}, 1996).

In accordance with their diverse structures, zinc finger proteins have been assigned multiple functions, including DNA recognition, transcriptional activation, RNA packaging, regulation of apoptosis, ubiquitination and many others (Coleman, 1992; Wolfe \textit{et al.}, 2000; Laity \textit{et al.}, 2001; Vazquez \textit{et al.}, 2007). More than 20 different zinc finger genes located on sex chromosomes or autosomes have been proposed to play a regulatory role in mammalian spermatogenesis (Noce \textit{et al.}, 1992; Pieler and Bellefroid, 1994; Yan \textit{et al.}, 2002).

The human \textit{ZNF230}, which maps to the short arm of chromosome 11 (11p15), encodes a C3HC4-type zinc finger protein motif (ring finger motif), and, consistent with a role in premeiotic or postmeiotic sperm development, one of its transcripts has been identified in abundance in the testicular tissue of fertile men, but neither in fetus nor in azoospermic patients. This suggested that \textit{ZNF230} may be involved in spermatogenesis, and loss of its expression may lead to azoospermia (Zhang \textit{et al.}, 2001). But so far, no clear biological function and mechanism had been elucidated.

In order to analyse the function of this novel zinc finger gene during vertebrate development, we decided to identify a potential orthologue in the teleost fish \textit{Danio rerio}. This animal has become widely used as a genetic model to uncover specific functions of unknown proteins (Dooley and Zon, 2000; Rubinstein, 2003). Being transparent early embryonic stages, easy to manipulate and highly reproductive makes the zebrafish an ideal animal system for molecular studies (Moro \textit{et al.}, 2007).

We here report the cloning and characterization of an 816 bp cDNA sequence, named \textit{rfn141}, which represents a candidate zebrafish orthologue of the human \textit{ZNF230} gene. By using whole-mount \textit{in situ} hybridization, RT-PCR, gene knockdown and overexpression analysis, we further showed its spatiotemporal expression pattern during early developmental stages.
Materials and Methods

Zebrafish embryo maintenance

Zebrafish AB strain was provided by National Zebrafish Resources of China, and maintained under standard laboratory conditions at 28.5 °C (Westerfield, 1993). Embryonic stages were identified by morphological features (Kimmel et al., 1995), and embryos in developmental stages of interest were fixed in 4% paraformaldehyde.

RNA extraction and reverse transcription

Total RNAs was isolated from adult zebrafish tissues using RNeasy Mini Kit (Qiagen) according to the manufacturer’s instructions. SuperScript TM Reverse Transcriptase using RNeasy Mini Kit (Qiagen) according to the manufacturer’s instructions. 

Cloning of the zebrafish rnf141 gene

Primers for 5’-RACE (366L 5’-ACGAGACGC CTCTACCATTCCATCC-3’) and the other three pairs of primers (70U 5’-CTTCCATTGGAGCCAAAGATGGGC C-3’ and 763L 5’- TAGATTITTTAAGGTCTGTGGG TG-3’; 338U 5’-AGGAGATGGAATGGTAGAGGCGT C-3’ and 619L 5’-GGGCTCTGGCCGCTCCACTTGTCA AT-3’; and 432U 5’-TAGTTCAAATGTGGCGGCAG AGGGA-3’ and 763L 5’- ATATATAGGTGTCTTT ATGGGGA-3’) were designed to obtain the complete coding sequence, based on the potential orthologue of ZNF230 in zebrafish. This orthologue sequence was acquired by reverse transcription, using cDNA from total RNAs of zebrafish tissue. The PCR products, including 5’- RACE products, were ligated into vector pGEM-T Easy Vector (Promega), cloned and sequenced bidirectionally. Sequences obtained by 5’-RACE and the other three fragments were assembled, and the contig was queried to the zebrafish genome database to determine its chromosomal location, and analyse its genomic structure. The deduced amino acid sequence was searched against InterPro Database for possible functional domains.

Multi-tissue RT-PCR

To reveal the tissue distribution and expression of zebrafish rnf141 gene, total RNA was extracted from embryos of various developmental stages (Kimmel et al., 1995) and several tissues of adult zebrafish. The gene-specific primers 488U 5’-GGATGGGACGGTAAAC AGTTGA-3’ (forward) and 683L 5’-GATCCGACATG ACCCAGGATTAG GC-3’ (reverse) were designed to amplify a 196 bp fragment of rnf141. Amplification was performed in 30 cycles as follows: 30 s denaturation at 94 °C, 30 s primer annealing at 62 °C and 1 min extension at 72 °C. The PCR products were electrophoresed on 1% agarose gel in 1 x TAE buffer and ethidium bromide stained.

Primers sequences used for amplifying 470 bp β-actin were 5’-TGTGGCCCTTGGACTTGTGACG-3’ (forward) and 5’-TAGAAAGCATTGGCGTTGGAGCA-3’ (reverse), according to Kaslin et al., (2004). In negative controls, ddH2O was used instead of cDNA template. The gene-specific primers were selected from two exons separated by an intronic sequence to identify possible amplicons from contaminating genomic DNA. All synthetic oligonucleotides were purchased from Invitrogen Corporation (CA, USA).

rnf141 gene knockdown and overexpression experiments

rnf141 morpholino antisense oligonucleotide (rnf141-MO, 5’-CCAGAAAGCTGCTGGCCCACCTTTG G-3’) was used to target rnf141 mRNA, and 5’-mpaired control morpholino (5’-CCACAAAACCTCCTGCACCATG TGG-3’) served as a control. Both were designed by using Gene-tools (Philomath, OR). The coding region of rnf141 was ligated into vector pcDNA3 (Invitrogen) and linearized by appropriate restriction enzymes for mRNA was synthesized by using mMESSAGE mMACHINE® Kit (Ambion). rnf141-MO and control-MO were injected into 1-2 cell zebrafish embryos by using a Model PLI-90 Pico-Injector.

Whole-mount in situ hybridization

rnf141 sense and antisense RNA probes were labeled with digoxigenin-11-UTP and synthesized by using DIG RNA Labeling Kit (SP6/T7) (Roche). The template for the probe was the entire 668 bp long cDNA. Whole-mount in situ hybridizations was performed as described by Westerfield (1995). Images were captured using an Olympus digital camera.

Results and Discussion

Zebrafish rnf141 has a C3HC4 zinc finger domain

We cloned zebrafish rnf141 based on the amino acid sequence of human ZNF230 and mouse znf230 by PSI-BLAST alignment and RT-PCR including RACE. As a result, four fragments were obtained that formed a 816 bp cDNA contig. This was submitted to GenBank (accession numbe BC071534) that encodes the same protein as 5’-TGTGGCCCTTGGACTTGTGACG-3’ (forward) and 5’-TAGAAAGCATTGGCGTTGGAGCA-3’ (reverse), according to Kaslin et al., (2004). In negative controls, ddH2O was used instead of cDNA template. The gene-specific primers were selected from two exons separated by an intronic sequence to identify possible amplicons from contaminating genomic DNA. All synthetic oligonucleotides were purchased from Invitrogen Corporation (CA, USA).

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The predicted open reading frame from 88 to 756 is 668 bp in length and encodes a polypeptide of 222 amino acid residues with a C3HC4 zinc finger domain from 146 to 266 amino acid residues.
**Expression analysis of rnf141**

To analyse the spatiotemporal expression of *rnf141* during early embryonic development, whole-mount *in situ* hybridizations were performed on two-cell stage to five-day-old embryos using an antisense probe. As a result, *rnf141* transcripts were already detected at the two-cell stage (Figure 2A), thus suggesting a maternal origin of the transcript. From the sphere stage (4 hpf) to the tail bud stage (10 hpf) (Figure 2C-E), the *rnf141* transcripts have a broad distribution. However, at the 5-somite stage (11.6 hpf), a characteristic pattern was displayed with marked staining in the notochord (Figure 2F), and at the Prim-5 stage of the pharyngula period, this pattern was displayed in the midbrain and hindbrain (Figure 2G). Following the long-pec stage (48 hpf), restricted signal localization was evident in the otic capsule, 4th ventricle, epiphysis and cerebellum (Figure 2H-I). When embryos reached the protruding-mouth stage (72 hpf), obvious signals were detected in the oral cavity and otic capsule (Figure 2J-K). In 5 dpf (120 hpf) embryos, an extensive *rnf141* expression was visible in the gut, with a restricted localization in the swim bladder (Figure 2L). To assess the specificity of the antisense probe, a sense probe was used in a parallel control experiment at all stages. With this sense probe no staining was detected in any embryo (Figure 3).

The consistency of hybridization experiments was confirmed by RT-PCR expression analysis performed on cDNAs from whole zebrafish embryos at various early developmental stages (Figure 2M).

Since the 1 kb transcript of human ZNF230 is only expressed in fertile male testis, whereas another 4.4 kb transcript was detected in many tissues; include heart, brain, skeletal muscle, kidney and pancreas (Zhang et al., 2001), we further addressed the question as to whether zebrafish *rnf141* maintains its ubiquitous spatial expression in adult stages. As shown in Figure 2N, RT-PCR based analysis demonstrated that almost all analysed tissues of adult fish do display a high content of *rnf141* transcripts.

In conclusion, these results of whole-mount *in situ* hybridization and RT-PCR analyses performed both on zebrafish embryos and adult tissues provide evidence that *rnf141* may have multiple functions. The detection of its transcripts in the CNS of early embryos, especially restricted in the notochord at the 5-somite of the segmentation period, suggests a function for *rnf141* in zebrafish development. Further analysis is ongoing in order to improve knowledge on the role of *rnf141*.

**rfn141 may play a part in normal dorsoventral patterning of zebrafish embryos**

To further study the potential function of *rnf141*, we first injected zebrafish embryos with synthetic *rnf141* mRNA. Injection of 200 pg *rnf141* mRNA caused 84% (n = 92) of the embryos to show phenotypes that are characteristic of embryonic ventralization at 24 hpf (Figure 4B). The expression of the shield-specific gene *goosecoid* was decreased at the shield stage (Figure 5A-c’). In contrast, the ventral markers *bmp2b* and *vent* expanded dorsally during gastrulation (Figure 5A-c’). The ratios of embryos with altered marker gene expression are summarized in Figure 5B.

To investigate the role of endogenous *rnf141*, a morpholino antisense oligonucleotide (*rnf141*-MO) was injected into one-cell embryos. As a result, 81% (n = 88) of the embryos injected with 12 ng *rnf141*-MO exhibited dorsIALIZED phenotypes at 24 hpf: complete loss of the yolk sac extension and partial loss of the caudal ventral fin (Figure 4C). The effects of *rnf141* knockdown on the expression of the marker genes *bmp2b*, *vent* and *goosecoid* (Figure 5Ab-b”) tended to be opposite to those of *rnf141* overexpression. In contrast, injection with 15 ng of control morpholino, which differs from *rnf141*-MO in five mismatched nucleotides, did not cause developmental defects (Figure 4E).

To test the efficiency of the morpholinos, fertilized eggs were injected with 12 ng *rnf141*-MO in combination with 100 pg of *prn141*-GFP DNA, an expression construct

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**Figure 1** - Nucleotide and predicted amino acid sequence of *rnf141* gene. The coding region (nucleotides 88-756) is in uppercase letters. The translation initiation codon is underlined. The stop codon at the 3’-end of the sequence is underlined and shaded. The deduced amino acid sequence (222 amino acids) is also shown below the nucleotide sequence. The predicted Ring-finger motif is shaded.
containing the full coding region of *rnf141* cDNA fused in-frame to a GFP coding sequence. At this dose of *rnf141*-MO, the injected embryos almost lacked green fluorescence from the GFP fusion protein (Figure 5I), while the

**Figure 2** - Expression analysis of *rnf141* in early embryos and adult zebrafish tissues. *rnf141* mRNA was initially detected at the 2-cell stage (A) and 4-cell stage (B), the signal become weaker at the sphere stage (C), shield stage (D) and bud stage (E). At the 5-somite stage (11.6 hpf), a characteristic pattern was displayed with marked staining in the notochord (F). At the 5-prim stage (24 hpf), a strong signal was detected in the head, particularly in the midbrain, hindbrain and the otic capsule (G). At the long-pee stage (48 hpf) the signal localization became restricted to the otic capsule, the 4th ventricle, as well as the epiphysis and tegmentum (H-lateral view from left; I-dorsal view). An even more restricted expressions was detected in the oral cavity and a restricted localization in swim bladder was found in 5 dpf embryo (L). Expression analysis of *rnf141* detected by RT-PCR in different developmental stage embryos (M) and adult zebrafish tissues (N). Abbreviations: mb, midbrain; hb, hindbrain; ep, epiphysis; cb, cerebellum; 4v, 4th ventricle; oc, otic capsule; ocv, oral cavity; sb, swim bladder.

**Figure 3** - Whole-mount in situ hybridization with *rnf141* antisense and sense probe. Hybridizations were performed on two-cell stage to five-day-old embryos using *rnf141* antisense probe (A) and sense probe (B).
same dose of rnf141-cMO injected embryos retained visible fluorescence (Figure 5H), suggesting that rnf141-MO could effectively block translation of rnf141 mRNA.

To test the specificity of rnf141-MO, a 5 mis-pair rnf141 mRNA corresponding to 5 mis-pair control morpholino was synthesized for rescuing the phenotype mediated with rnf141-MO. These results showed that the rnf141-MO-induced dorsalization could be neutralized by coinjection with a smaller amount of 5 mis-pair rnf141 (Figure 4D), suggesting that rnf141-MO specifically targets rnf141.

In conclusion, knockdown of rnf141 by using special morpholino-induced abnormal outcomes, including inordinate development of the CNS with an atrophic hindbrain, thin and crooked notochord, as well as disappearance of yolk sac extension, abnormality of axis, and partial loss of the caudal ventral fin. These embryos are characteristic of weakly dorsalized phenotypes, reminiscent of mini fin (mfn) and lost-a-fin (laf) mutant embryos, which were first described by Mullins et al (1996) and were subsequently found to be caused by inefficient BMP signaling (Bauer et al., 2001; Connors et al., 1999; Mintzer et al., 2001). Overexpressing this gene by injection of rnf141 mRNA may caused embryos to show ventralized phenotypes. We also noted that rnf141-MO-induced dorsalization could be neutralized by coinjection of a smaller amount of rnf141 mRNA, suggesting that rnf141-MO specifically targets rnf141 mRNA. The expression of ventral markers (bmp2b and vent) and of a dorsal marker (goosecoid) were impacted by altered expression of rnf141, thus suggesting that zebrafish rnf141 may participate in normal dorsoventral embryonic patterning. Further research is needed to better understand the respective biological pathway(s) and improve the knowledge on the function of rnf141.
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