Epidermis-restricted expression of zebrafish cytokeratin II is controlled by a $-141/+85$ minimal promoter, and cassette $-141/-111$ is essential for driving the tissue specificity

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Abstract  We isolated a 2.3 kb DNA segment from the upstream region of the zebrafish cytokeratin II (zfCKII) gene. Transgenic embryos, produced by using a series of 5’ deletions linked to the red fluorescent protein (RFP) reporter, showed that the $-141/+85$ segment of zfCKII directed RFP expression in epidermal cells, whereas the $-111/+85$ segment did not. When $-141/-111$ was deleted from $-355/+85$ and microinjected into one-celled embryos, no fluorescence was observed at later stages, indicating that the $-141/-111$ segment is required for green fluorescent protein expression in epidermal cells. Furthermore, when a putative KLF-binding site at $-119/-117$ was mutated, RFP expression rates and intensities were reduced dramatically, although still observed, suggesting that $-119/-117$ within $-141/-111$ is a key cis-element for controlling epidermis-specific expression of the zfCKII gene. Finally, we generated a zebrafish transgenic line, Tg(zfCKII(2.3):RFP), which carries an upstream 2.3 kb regulatory region of the zfCKII gene fused with RFP. The expression pattern in the epidermal cells of Tg(zfCKII(2.3):RFP) fish recapitulated that of the endogenous gene. F2 embryos derived from Tg(zfCKII(2.3):RFP) males crossed with wild-type females revealed that the earliest onset of RFP expression was at the sphere stage, indicating that this transgenic approach can be used for studying zygotic expression of maternally inherited genes.

Key words  cytokeratin  epidermal cells  germline transmission  red fluorescent protein  transgenic  zebrafish

Introduction

There are many families of cytoplasmic filament proteins. Keratin is one such family, having intermediate size (10 nm) and expression specifically in epithelial cells as cytoskeletal proteins. At least 30 related members in the cytokeratin family are known and are encoded by complex multiple genes. Type I keratins (K9–K20) are more acidic ($pI = 4–6$), whereas type II keratins (K1–K8) are neutral or basic ($pI = 6–8$). Unlike other intermediate filament (IF) proteins, cytokeratins form obligatory heteropolymers, consisting of equal numbers of type I and type II keratins (Fuchs and Weber, 1994; Klymkowsky, 1995; Rogers et al., 2004) as specific pairs, and form filaments by the coiled-coil interaction (Hatzfeld et al., 1987).

There are 19 human keratin isoforms (K1–K19), and specific pairs of keratin isoforms determine tissue identity and differentiation state. For example, human keratins K5 (type II) and K14 (type I) together form the extensive IF network of mitotically active basal cells in all stratified epithelia (Byrne et al., 1994). Expression of human keratin is restricted: E keratins are found in stratified epithelium (including epidermis), and S kerat-
tins are found in simple epithelium. However, in rainbow trout, both E and S keratins are expressed in intestinal simple epithelium (Markl and Franke, 1988). In zebrafish, S keratins are expressed both in stratified epithelium (including epidermis) and simple epithelium (Imboden et al., 1997). In addition, fish keratin can be found in mesenchyme-derived cells and certain nerve cells (Giordano et al., 1989, 1990), but mesenchymal tissue usually expresses vimentin in humans. This evidence indicates that fish epidermal keratin diversified independently from that of mammalian epidermal keratin. Thus, the mechanism of transcriptional regulation of the keratin in fishes should be studied.

In mammals, the regulation of keratin genes by a variety of cis-acting elements has been studied. For example, a keratinocyte-specific enhancer (the 10 bp 5’-ACCTGCAGGC-3’) has been identified in the K14 gene (Vassar et al., 1989; Leask et al., 1990). Nuclear proteins, such as GKL/KLF4 and Sp1, which interact with the upstream region of the human K19 keratin gene, have also been identified (Brembeck and Rustgi, 2000). Rhodes and Oshima (1998) reported that the expression of human and mouse K18 is activated by the transcription factors c-jun and c-fos, respectively, by means of an outer AP1 site in the first intron. Additionally, Popa et al. (2004) reported that the AP-2 transcription factor family is presumed to play an important role in the regulation of the keratinocyte squamous differentiation program. There is extensive knowledge about the transcriptional regulation of keratin genes in mammals; however, the mechanism of transcriptional regulation of keratin in fish has not been reported to date, although fish cytokeratin has been characterized from rainbow trout (Markl and Franke, 1988; Markl et al., 1989; Schaffeld et al., 2002a, 2002b), goldfish (Druger et al., 1992), catfish (Ainis et al., 1995), lamprey (Zacone et al., 1995), mosquito fish (Arenas et al., 1995), common carp (Groff et al., 1997), zebrafish (Conrad et al., 1998; Chua and Lim, 2000; Schaffeld et al., 2003; Wang et al., 2006b), and shark (Schaffeld et al., 1998).

Previously, a zebrafish type II cytokeratin (zfCKII) cDNA was isolated and characterized (Chua and Lim, 2000). zfCKII was expressed uniformly in the skin, fins, scale epidermis, retina, cornea, neurons, and glial cells of the brain and spinal cord as well as in chondrocytes of the skull of the adult fish. The tissue-specific expression of zfCKII is of particular interest because of the broad diversity of zfCKII-positive epithelial cells. In the present study, a 2.3 kb zfCKII promoter was isolated and its function was assayed in both transient and stable transgenic lines. In stable transgenic lines, red fluorescent protein (RFP) expression faithfully reflected the expression of the endogenous zfCKII gene. Microinjections of serial deletions of the upstream segment of zfCKII fused with RFP were performed. We found that a cis-element of the DNA segment in the zfCKII gene is required for epidermal cell-specific expression of zfCKII and that a minimal region upstream of the transcription start site (141 bp) is required for this expression restricted to epidermal cells.

Materials and methods

Experimental fish

Zebrafish AB strains (wild-type, wt) were kept under a 14 hr light and 10 hr dark photoperiod at approximately 28.5°C. After fertilization, the eggs were collected and cultured in an aquarium. Embryonic cleavage and somite formation were observed with a light microscope to determine the developmental stages (Kimmel et al., 1995).

Isolation of the zfCKII upstream regulatory region

The 5’-flanking region of the zfCKII gene was isolated according to the instructions for the Universal GenomeWalker Kit (BD Biosciences Palo Alto, CA). Briefly, DNA was extracted from zebrafish embryos (Westerfield, 1995), digested with restriction enzymes, and ligated to a GenomeWalker adaptor to produce GenomeWalker libraries. DNA fragments containing the putative upstream region were isolated after two successive polymerase chain reaction (PCR)-based DNA walking in GenomeWalker libraries. The primary PCR and the outer adapter primer (AP-1) products were obtained using the GeneAmp PCR System with five cycles of 94°C for 5 sec and 72°C for 3 min, followed by 30 cycles of 94°C for 5 sec and 68°C for 3 min, and a final extension at 68°C for 7 min. For each round of genomic walking, the diluted primary PCR products served as templates for the secondary nested PCR with a nested gene-specific primer (CKIIIR) and the nested adaptor primer (AP-2). The secondary PCR products were analyzed and purified from agarose gels and cloned into pGEMT easy Vector (Promega, Madison, WI) to produce pT7CKII. The primers used in the genomic walking and subsequent cloning are listed in Table 1.

5’-Rapid amplification of cDNA end (5’-RACE)

The protocols of 5’-RACE are followed by the manufacturer’s instructions (FirstChoice™ RLM-RACE; Ambion, Austin, TX) with minor revision. Briefly, first-strand cDNA used for 5’-RACE was performed as described before (Chen et al., 2001), and then using 5’-RACE outer primer and CKII-271R for the first PCR, and primers CKII-177F and 5’-RACE inner primer for the nesting. Amplified DNA fragments were sub-cloned and sequenced as described above.

Construction of chimeric reporter gene fusions

The upstream segments of zfCKII and of the serial deletion derivatives were fused with the RFP reporter gene by cloning the various 5’ deletion fragments into the polylinker region of the vector pDsRed2.1 (Clontech, Palo Alto, CA) upstream of a promoterless RFP gene. To generate pCKII2.3K-RFP, pT7CKII was digested by EcoRI and a 2.3 kb EcoRI-digested fragment was purified and ligated to EcoRI-digested pDsRed2.1 (Clontech). The deletion fragments of 1.7 kb, 1.2 kb, 526 bp, 440 bp, 228 bp, 194 bp, and 164 bp corresponded to the CKII gene from −1790 to +85 (−1709/+85), −1618/+85, −441/+85, −355/+85, −141/+85, −110/+85, and −80/+85, respectively. They were amplified from pT7CKII using forward primers CKII(−1.7K)F, CKII(−1K)F, CKII(−441)F, CKII(−355)F, CKII(−141)F, CKII(−110)F, and CKII(−80)F, respectively, and the common reverse primer CKIIIR. These resulting PCR products were cloned into pGEM-T Easy Vector (Promega) to produce pT7CKII1.7K, pT7CKII1K, pT7CKII(−441/+85), pT7CKII(−355/+85), pT7CKII(−141/+85), pT7CKII(−110/+85), and pT7CKII(−80/+85), respectively. Then, all plasmids were digested by EcoRI and ligated to EcoRI-digested pDsRed2.1 (Clontech) vectors to generate pCKIII.
Table 1 Oligonucleotides used in this study

<table>
<thead>
<tr>
<th>Symbols</th>
<th>Nucleotide sequences (5’–3’)</th>
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<tbody>
<tr>
<td>5’RACE outer</td>
<td>GCTGATGGGGATGAATGAACACTG</td>
</tr>
<tr>
<td>CKII–271 R</td>
<td>CAAATCTCTCCACCTGATGG</td>
</tr>
<tr>
<td>5’RACE inner</td>
<td>CGCCTGACGACACTGTTTGGGCTGGTTTGTGCTGGATG</td>
</tr>
<tr>
<td>CKII–117 R</td>
<td>GTGCTGCCATTGACGCTGTCGTTAGG</td>
</tr>
<tr>
<td>AP-1</td>
<td>GTAATACGACTCATTAGGGC</td>
</tr>
<tr>
<td>AP-2</td>
<td>ACTATAGGGCAAGCGGTTG</td>
</tr>
<tr>
<td>CKII (1.7K) F</td>
<td>CTTGGAACATGGTATTGTAGTATTGCTCTC</td>
</tr>
<tr>
<td>CKII (1.10 K) F</td>
<td>GACTGTCAAGGACATTTAAAAATCGAC</td>
</tr>
<tr>
<td>CKII (1.10 K) Fm</td>
<td>GACCAACGGATACAAATGGGAC</td>
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<tr>
<td>CKII (1.11 K) F</td>
<td>GCACCTAAGACAACTGAGGGCCC</td>
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<tr>
<td>CKII (1.11 K) Fm</td>
<td>CTGGTATGTAAATAAGAGGGGAC</td>
</tr>
<tr>
<td>CKII (1.11 K) R</td>
<td>GCAGGCGAGAGCGATAAGGCCC</td>
</tr>
<tr>
<td>CKII (80) F</td>
<td>TTGGCCAGTGAGCCCTCC</td>
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<tr>
<td>CKII del (117/111) F</td>
<td>CAGCAAATGGGACAGACAGACTGAGAAGAAGGG</td>
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<tr>
<td>CKII del (117/111) Fm</td>
<td>CAGGGCTCTTGCTGCAATTTGGACGCTGGTATGGG</td>
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<tr>
<td>CKII (117/111) R</td>
<td>AATTTCTTTTTCTTCGTTCAAGATGATACACACGCC</td>
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7K-RFP, pCKII K-RFP, pCKII (144/85)-RFP, pCKII (355/85)-RFP, pCKII (110/85)-RFP, and pCKII (80/85)-RFP, respectively. All constructs were confirmed after sequencing.

To generate pCKII (560/85)-RFP, a pT7CKII was digested by EcoRI and HindIII, and a 644 bp fragment was ligated to EcoRI–HindIII-digested pDsRed2.1 (Clontech). To generate pCKII (355/85)-RFP, which contains a fragment spanning 355/85 but with 141/111 deleted, two PCR products were obtained by amplifying the pT7CKII template using primers of CKII del (117/111) F and CKII. After annealing, the PCR products served as templates for amplification of primers CKII (355/355) F and CKII. To generate plasmid pCKII (141/85)KLFm-RFP, which contains a fragment spanning 141/111 but containing a mutated KLF-binding site (−182GGG to −192AAA), the two PCR products were annealed after they were amplified from the pT7CKII template using primers of CKII (117/111) F and CKII (−141/−110) Fm and CKII. After annealing, the PCR products served as templates for amplification of the mutated segment by primers CKII (117/111) F and CKII. These resulting PCR products were cloned into pGEM-T Easy Vector (Promega) to produce pT7CKII (355/85)Δ (141/111) and pT7CKII (141/85)KLFm, respectively. Then, pT7CKII (355/85)Δ (141/111) and pT7CKII (141/85)KLFm were digested by EcoRI and ligated to EcoRI-digested pDsRed2.1 (Clontech) vectors to generate pCKII (355/85)Δ (141/−111) RFP and pCKII (141/85)KLFm, respectively. To construct plasmid pEGFPm (141/−111), which contained the −141/−111 region of CKII fused with a cytomegalovirus (CMV) minimal TATA promoter and with the enhanced green fluorescent protein (EGFP) reporter gene, primers of CKII (141/−111) F and CKII (141/−111) R were used and the PCR product was inserted into the EcoRI-digested pEGFPm (Chen et al., 2003).

Preparation of plasmids for microinjection

Plasmids pCKII 2.3K-RFP, pCKII 7.4K-RFP, pCKII 1K-RFP, pCKII (560/85)-RFP, pCKII (355/85)-RFP, pCKII (141/85)-RFP, and pCKII (110/85)-RFP, were linearized by XhoI and recovered from a 0.8% low-melting temperature gel (FMC BioProducts, Philadelphia, PA). DNA intensity and gel absorbancy were measured with a GeneQuant II calculator (Pharmacia Biotech, Hong Kong, China). DNA samples were resuspended at a concentration of 25 ng/μl in double-distilled water mixed with 0.1% (v/v) phenol red before use.

To get a quantitative view of the relative strength of the promoter constructs, we classified RFP-expressing embryos into three levels based on the expression pattern in the skin epidermis: strong, medium, and weak. The strong level included embryos displaying RFP expression in approximately 90% of skin epidermis throughout the body; the medium level included embryos showing RFP expression in 20%–90% of skin epidermis; and the weak level included embryos having RFP-positive signal in less than 20% of the skin epidermis.

Generation of transgenic germine transmitted zebrafish

The 2.3 kb pCKII 2.3K-RFP was linearized with XhoI and resuspended in 0.1 M KCl to a final concentration of 50–100 ng/μl with 0.2% phenol red as a tracer. The DNA solution was microinjected into the cytoplasm of one-cell stage embryos. After microinjection, red fluorescent signal was observed in 24 hr post-fertilization (hpf) embryos. All RFP-positive embryos were raised to adulthood. Pairs of transgenic founder (F0) fish were crossed with wt zebrafish to generate a heterozygotic F2 generation. After screening, RFP-positive F1 embryos were raised to adulthood and crossed with wt zebrafish to generate a heterozygotic F2 generation.

Embedding and cryosectioning

The embedding and cryosectioning protocols followed were those of Chen and Tsai (2002), with minor modifications. In brief, embryos were fixed with 4% paraformaldehyde for 4 hr at 25°C, dechorionized, mounted with 5% sucrose containing 1.5% agarose for 1 hr, cut into cubes approximately 5 × 5 × 5 mm, and stored in...
5% sucrose at 4°C. All embryos were embedded using Cryomatrix (Shandon, Waltham, MA), and cryosectioning (12 μm thick) was performed with a Microm Cryosector (Heidelberg, Germany).

Microscopy

Transgenic embryos were observed hourly, especially from 1 to 48 hpf, under a stereo dissecting microscope (MZ12, Leica, Wetzlar, Germany) equipped with a fluorescent module having a GFP or DsRed filter cube (Kramer Scientific, Hampton, NJ). Photographs were taken with an S2 Pro digital camera (Fuji, Tokyo, Japan) when embryos developed at specific stages.

Results

Upstream regulatory region of the zebrafish CKII gene

Previous studies have shown that zfCKII mRNA is expressed uniformly in the skin, fins, scale epidermis, retina, cornea, neurons, and glial cells of the brain and spinal cord as well as in chondrocytes of the skull of the adult fish (Chua and Lim, 2000). To investigate the molecular mechanism of zfCKII gene expression in epidermal cells, a 2.3 kb DNA segment of the upstream region of zfCKII was isolated and its partial DNA sequence was determined (Fig. 1; the sequence (2219/441), GenBank Accession No. DQ144236). We also performed a 5’-RACE experiment to determine the transcription start site and the length of the 5’-untranslated region of ZFCKII gene. As shown in Fig. 1, the numbers indicate nucleotide positions with the transcription start site as +1. Using a string-based search query, the TESS program (http://www.cbil.upenn.edu/tess/) revealed that the putative binding sites of the transcription factors within the proximal (2219/441) segment were the AP1 site (CATGAATCATA, (2219/141)), the Oct-1 site (TGGTTTGCAT, (2219/70)), two Sp1 sites (GGGTGTGGC, −196/−188; and GGGATTGGGCC, −85/−75), and a TATA box (TATAAAA, −46/−40; Fig. 1).

Transient expression of RFP is driven by the upstream 2.3 kb segment of the zfCKII gene in zebrafish embryos

The tissue-specific expression mediated by the isolated upstream segment of zfCKII was investigated using a transient transgenic approach. The earliest RFP expression in the embryos microinjected with pCKII2.3K-RFP was in the head region at 19 hpf (data not shown). At 3 days post-fertilization (dpf), the RFP signals were extended to the body, including the head (Fig. 2A), skin epidermis, pectoral fins (Fig. 2B), and caudal fins (Fig. 2C). At 14 dpf, the red fluorescence was observed in the anal fins and in the retina (Figs. 2D, 2F). These transient expression studies indicated that the upstream segment in the zfCKII2.3K contained regulatory elements that drove RFP expression in a tissue-specific manner.

Functional analysis of the zfCKII regulatory sequence

To determine whether the cis-acting element is sufficient and required for the epidermis-specific expression of zfCKII, serial deletions of an upstream region of zfCKII were generated and fused with RFP cDNA. These constructs were microinjected into zebrafish fertilized eggs. Embryos injected with DNA fragments containing the −2219, −1709, −1068, −560, −441, −355, and −141 bp upstream sequences [pCKII2.3K-, pCKII1.7K-, pCKII1K-, pCKII(−560/+85)-, pCKII (−441/+85)-, pCKII(−355/+85)-, and pCKII(−141/+85)-RFP, respectively] displayed high RFP expression rates (63%–85%, Table 2) and 100% RFP-positive
signals in the skin epidermis at 3 dpf (Fig. 3A and Table 2). To obtain a quantitative view of the relative strength of the promoter constructs, we classified RFP-expressing embryos into three levels based on the expression pattern in the skin epidermis: strong, medium, and weak with reference to the previous studies (Ju et al., 2003; Lin et al., 2004). The strong level included embryos displaying RFP expression in approximately 90% of the skin epidermis throughout the body; the medium level included embryos showing RFP expression in 20%–90% of skin epidermis; and the weak level included embryos having RFP-positive signal in less than 20% of the skin epidermis (Fig. 3C). As shown in Fig. 3B, embryos displayed relatively high RFP expression rates in strong (59%–78%) and medium (20%–38%) levels when the injected upstream sequences were longer than $141 \text{bp}$. In addition to red fluorescent skins, embryos injected with DNA fragments containing $1709, 1068, 560$, and $441 \text{bp}$ upstream sequences also displayed high expression rates (49%–68%) of RFP-positive signals in muscle (Table 2; Fig. 3C). However, the $pCKII(-141/+85)$-RFP- and $pCKII(-141/+85)$-RFP-injected embryos displayed red fluorescence in 1.5% and 3.2% of muscle fibers (Table 2; Fig. 3C).

It was found that the $141 \text{bp}$ upstream sequence was able to drive skin-restricted expression of zfCKII. On the other hand, embryos injected with DNA fragments containing less than 110 bp [$pCKII(-110/+85)$-RFP and $pCKII(-80/+85)$-RFP] did not give off RFP-positive signals (Fig. 3), even when the concentration of injected DNA was increased to 100 ng/µl (data not shown). On the basis of this evidence, we suggest that the proximal part of the $141/185$ sequence of zfCKII is a minimal cis-element for controlling specific expression.

Cassette $-141/-111$ was able to direct epidermal cell-specific expression

Interestingly, we found that, when a DNA fragment, in which $141/111$ was deleted from the zfCKII $-355/+85$ [$pCKII(-355/+85)\Delta(-141/-111)$-RFP], was injected into one-celled embryos, only 1% (two of 182, Table 2) of surviving embryos displayed red fluorescence. This result suggests that the $141/-111$ cassette can drive skin-restricted expression.
rescence in muscle, and none of the 182 embryos displayed RFP-positive skin epidermis (Table 2). To determine whether cassette $-141/-111$ was able to direct epidermal cell-specific expression of the reporter gene, we used plasmid pEGFPm($-141/-111$), in which a mini-promoter of CMV (TATA box only) was fused with GFP and one copy of the $-141/-111$ cassette (Fig. 4A). Only 2% of the 136 embryos injected with pEGFPm were GFP-positive, and none had signal detected in the skin (Fig. 4B, left panel; Table 2). However, the expression rate of epidermal cell-specificity in the transgenic embryos injected with pEGFPm($-141/-111$) was 45% (Fig. 4B, right panel; Table 2). Furthermore, a mutation plasmid, pCKII$(-141/+85)$KLFm-RFP, in which $-119$GGG$-117$ was mutated to $-119$AAA$-117$, was injected into the embryos and revealed a 29% RFP expression rate (Table 2). Interestingly, most of the pCKII$(-141/+85)$KLFm-RFP-injected embryos displayed medium (43%) and weak (52%) RFP expression, and only 5% displayed strong RFP expression (Fig. 3B). Based on these observations, we propose that cassette $-141/-111$ was able to direct epidermal cell-specific expression and that $-119$GGG$-117$ is an important element within cassette $-141/-111$.

RFP expression in germline zfCKII-RFP transgenic zebrafish

To generate stable transgenic lines for further analysis of the expression mechanism of zfCKII, embryos injected with pCKII2.3K-RFP and exhibiting RFP expression were collected and raised to adulthood. Founder fishes (F0) were crossed with wt or crossed with each other to generate F1 embryos. Of 63 founder fish tested, one line, Tg(zfCKII(2.3):GFP), produced embryos that expressed RFP in the epidermis, but 5% (11 positive of 234) of embryos expressed RFP at 24 hpf, suggesting that the transgenic founder is a germline mosaic, which is a common finding in transgenic fish at the founder generation (reviewed by Gong and Hew, 1995). The F2 inheritance rate of RFP-positive F1 individuals of the Tg(zfCKII(2.3):RFP) line is 48.57% (102 of 210 total embryos), indicating that there was a single insertion site of the transgene in the genome.

All F2 embryos (154 of 154 embryos) derived from an F1 female crossed with a wt male exhibited red fluorescent signal at one-cell (Fig. 5A), blastula (Fig. 5B), and segmentation (Fig. 5C) stages. Red fluorescence became stronger and extended to the body at 3 dpf (Fig. 5D). In contrast, none of the F2 embryos derived from an F1 male crossed with a wt female showed red fluorescence at the one-cell (Fig. 5E) and blastula stages (Fig. 5F), yet 46.9% of 145 embryos began to display red fluorescence at 14 hpf (data not shown). The red fluorescence became stronger at 24 hpf (Fig. 5G), and extended to the trunk in a non-uniform expression manner (Fig. 5H). This evidence strongly demonstrates that the zfCKII transcripts are maternally inherited.

Cryosectioning of 4 dpf embryos derived from the Tg(zfCKII(2.3):RFP) line revealed RFP signal distributed in the retina (Fig 5I), head epidermis (Fig. 5I), trunk epidermis (Fig. 5K), and pectoral fins (Fig. 5K). In addition, we think the faint red fluorescent signal in the brain in the 4 dpf larva section (Fig. 5I) represents real signals reflecting the endogenous gene expression. In the 1-month-old juveniles, red fluorescence was also observed on the body surface, on the cornea (Fig. 5M), and in the epidermal cells of the scales (Fig. 5N). This red fluorescence was found only in the outer region of the scale (Fig. 5O). From these observations, we conclude that Tg(zfCKII(2.3):RFP) fish recapitulate the endogenous zfCKII expression patterns.
Discussion

In this study, we isolated a 2.3 kb upstream regulatory sequence of the zfCKII gene. Analyses of transient as well as stable transgenic lines revealed that this 2.3 kb segment is able to recapitulate the endogenous zfCKII expression patterns. Deletion analyses were performed and showed that a proximal −141/+85 sequence of zfCKII is a minimal cis-element for controlling expression specificity. Finally, we identified a 31 bp (−141/−111) segment, which is able to drive GFP expression in the skin epidermis.

Many transcription factors (c-jun, c-fos, C/EBP, and SP-1) and their binding sites (AP1, CCAAT-, and GC-box) have been reported to play important roles in regulating mammal keratin genes at transcription (Brembeck and Rustgi, 2000; Sterneck et al., 2006; Wang et al., 2006). As shown in Fig. 1, AP1, CCAAT-, and GC-box exist in the −441/+85 segment of the zfCKII gene. We examined the putative transcription factor-binding sites in zebrafish k8 (now renamed as k4) and k18 genes (Gong et al., 2002; Wang et al., 2006) and found that AP1, CCAAT-, and GC-box are located in the upstream regions of k4 and k18. Thus, we propose that zebrafish keratin genes zfCKII, k4, and k18 shared some common transcriptional regulatory mechanisms with those of keratin genes in mammals.

Systematic analyses of mammal keratin revealed that the tissue specificity is most probably conferred by sequences close to the TATA box. But in some cases, the enhancers for directing tissue specificity are located at the first intron (Rhodes and Oshima, 1998) and the 3' flanking sequence (Hu and Gudas, 1994). A transgenic analysis of the human k5 promoter revealed that 90 bp of the 5' flanking sequences contain sufficient information to direct expression to keratinocytes (Byrne and Fuchs, 1993). Mahony et al. (2000) reported that a...
120 bp mouse keratin 6a mini-gene contains sufficient sequence information to direct uniform and tissue-specific expression. In zfCKII, we showed that the proximal –141/+85 sequence of zfCKII is a minimal cis-element and that a 31 bp (–141/–111) segment is capable of driving GFP expression in the skin epidermis. Sequence analysis of –141/–111 segment revealed that there is one KLF-binding site (CACGGGG–117). KLF has been reported to be involved in regulating tissue-dependent transcription of the keratin 19 gene (Brembeck and Rustgi, 2000). We have shown a mutation plasmid, pCKII (–141/+85)KLfm-RFP, in which –111CA–GGGG–117 has been mutated to –119AAA–117, resulting in an altered KLF-binding site. Comparing the pCKII (–141/+85)-RFP with the pCKII (–141/+85)KLfm-RFP-injected groups, we found that the RFP expression rates were significantly reduced (81% versus 29%, respectively; Table 2). Only 5% of pCKII (–141/+85)KLfm-RFP-injected embryos displayed strong RFP expression, a rate much lower that of the pCKII (–141/+85)-RFP-injected group (78%, Fig. 3B). These observations strongly suggest that –122CACGGGG–117 might be an enhancer for the zfCKII gene.

Embryos injected with pCKII2.3K-RFP displayed normal skin-specific expression (Table 2). However, embryos injected with pCKII1.7K-, pCKII1K-, pCKII (–560/+85)-, pCKII (–441/+85)-, pCKII (–355/+85)-, and pCKII (–141/+85)-RFP displayed ectopic expression in muscle (Table 2). Thus, we propose that there is a muscle repressor in the –2.3 to –1.7 kb region.

Chua and Lim (2000) showed that zfCKII mRNA was inherited maternally. The mechanism of transcriptional regulation of maternally inherited genes is difficult to study because of the interference of maternally produced mRNA. Here, for the first time, we demonstrate that an upstream 2.3 kb segment of the zfCKII gene is able to drive zfCKII expression before mid-blastula transition and is sufficient to recapitulate the endogenous zfCKII transcription. We used the Tg(zfCKII(2.3):RFP) line for studying the regulatory mechanism. F2 embryos derived from Tg(zfCKII(2.3):RFP) males crossed with wt females displayed red fluorescence without an interfering maternal effect. Thus, this fish line can be used as an excellent tool for studying zygotic expression of maternally inherited genes. In fact, we can knock-down several putative transcription factors, such as KLF, c-Jun/Fos, or SP1, by injecting the morpholinos into the F2 embryos that were produced by mating Tg(zfCKII(2.3):RFP) males with wt females to study the zygotic regulation of zfCKII. This transgenic line should provide new insights into zfCKII expression at the transcription level in early embryogenesis.

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