Identification and tissue distribution of mRNAs encoding salmon-type calcitonins-IV and -V in the rainbow trout

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Abstract

Four types of calcitonin are produced in salmonid fish, although their functional diversity is almost unknown. To explore the significance of these isoforms, we have characterized salmon-type calcitonin (sCT) mRNAs in the rainbow trout (Oncorhynchus mykiss), and examined their tissue distribution. In addition to the previously isolated sCT-I cDNAs, two new forms of sCT cDNA were cloned from the ultimobranchial gland, and one of them (sCT-IV cDNA) was predicted to encode an N-terminal peptide of 80 amino acid residues, a putative cleavage site Lys-Arg, sCT-IV, a cleavage and amidation sequence Gly-Lys-Lys-Arg, and a C-terminal peptide of 18 amino acids. The sCT-IV precursor was 78% identical with the rainbow trout sCT-I precursors. The other cloned cDNA encoded a precursor for a novel CT, sCT-V. The sCT-V peptide was different from sCT-IV by only one amino acid residue: Val at position 8 in the latter was replaced by Met. The sCT-V precursor had 80 and 90% identity with the sCT-I and -IV precursors respectively. No cDNA clones were obtained for sCTs-II or -III.

Tissue distribution of sCT-I, -IV and -V mRNAs was examined by RT-PCR and specific cleavage with restriction enzymes. An amplified fragment from sCT-I mRNA was detected not only in the ultimobranchial gland, but also in the gills, testis and ovary. RT-PCR analysis coupled to restriction digestion further revealed that sCT-IV mRNA was expressed in both the testis and the ultimobranchial gland. The expression sites of sCT-IV mRNA were localized to the Leydig cells of the testis and to the parenchymal cells of the ultimobranchial gland, by in situ hybridization histochemistry. Although the amino acid sequence of sCT-V peptide was nearly the same as that of sCT-IV, the sCT-V gene showed a much wider pattern of expression: the band amplified by RT-PCR was detected in all the tissues examined except the kidney, gills and blood cells. The sCT-V mRNA was shown to be localized in the parenchymal cells of the ultimobranchial gland, but not in other tissues at the cellular level, suggesting very low expression of sCT-V mRNA in those tissues. Our results show different patterns of tissue expression of three types of sCT genes in the rainbow trout, suggesting that sCTs-I, -IV and -V might differ in their local actions.

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Introduction

Calcitonin (CT) was initially discovered as a hypocalcaemic hormone in mammals. It is abundantly produced by the parafollicular cells of the thyroid gland, and principally inhibits osteoclast-mediated bone resorption (Sexton et al. 1999). It is also reported that CT is secreted by a variety of extrathyroidal tissues (Becker et al. 1979, Ding et al. 1994), and that CT receptor mRNA is expressed at many of these sites, such as the brain (Sheward et al. 1994), kidney, lung and stomach (Kuestner et al. 1994), suggesting diverse local actions of CT.

Down the phylogenetic lineages of vertebrates, non-mammalian calcitonin is produced by the ultimobranchial gland, and the sequences of CT peptides and mRNAs have been determined in a number of species, from fish to mammals (Sasayama 1999, Suzuki et al. 1999b). Sequence analysis reveals that CT is a 32 amino acid polypeptide that is derived from a larger precursor molecule, and further that some species produce
more than one form of CT (Suzuki et al. 1999b). Within the Salmonidae, four types of CT, sCTs-I to -IV, have been reported to date, and it has been suggested that pink salmon could express sCTs-I, -II and -IV (Jansz et al. 1996). Jansz et al. (1996) also suggested that sockeye and chum salmon possess both sCT-I and sCT-II, whereas coho salmon has sCTs-II and -III. It has also been suggested that, in fish, CT might control the uptake of dietary calcium from the digestive tract (Suzuki et al. 1999a), suppress osteoclastic activities in the scales (Suzuki et al. 2000a), and be involved in reproductive physiology (Bjornsson et al. 1986, Maubras et al. 1990). However, little is known about differences between the physiological roles of the isoforms of CT existing in the same species. Investigation of the tissue distribution of the isohormones is a prerequisite for the understanding of their functional differences. In the present study, we have cloned and sequenced cDNAs encoding two types of sCT (IV and V) from the ultimobranchial gland of rainbow trout, and compared the tissue distribution of their mRNAs with that of sCT-I mRNA.

Materials and methods

Animals and sampling

Two-year-old rainbow trout (Oncorhynchus mykiss) of both sexes were collected at Fuji Trout Hatchery, Shizuoka, Japan, in March and May. Ultimobranchial glands and other organs were carefully dissected out, and blood cells were collected by centrifuging the blood at 3000 g for 4 min. The samples were immediately frozen in liquid nitrogen, and stored at -80 °C in a deep-freezer. In addition, various tissues of adult trout were fixed in 4% paraformaldehyde at 4 °C overnight for histological analysis. Adult rainbow trout of a different strain were obtained from a commercial dealer, Yabe Trout Hatchery, Tokyo, Japan, in January. Wistar rats of 6 weeks of age (Japan SLC, Shizuoka, Japan) were anaesthetized with ether, and dissected to provide tissues for RT-PCR analysis.

Construction of cDNA library and DNA cloning

Total RNA was extracted from 120 ultimobranchial glands of the trout using TRIzol reagent (Gibco BRL, Rockville, MD, USA), and poly(A)+ RNA was prepared with oligo-dT-coated latex beads (Oligotex-dT30 super; Takara, Kyoto, Japan), according to the manufacturer’s instructions. cDNA was prepared with a ZAP Express cDNA synthesis kit (Stratagene, La Jolla, CA, USA), and then a cDNA library was constructed using a ZAP Express cDNA Gigapack III Gold cloning kit (Stratagene). The resultant cDNA library contained 7·0 × 10⁶ recombinant lambda phages, and was amplified to 7·6 × 10⁸ plaque-forming units (pfu)/ml.

Preparation of cDNA probe

Poly(A)+ RNA (0·5 µg) from the ultimobranchial gland was reverse-transcribed in 20 µl buffer containing 80 pmol oligo-dT19 primer, 1 mM dNTP, 20 units RNase inhibitor (Toyobo, Osaka, Japan), and 10 units Rous-associated virus 2 (RAV-2) reverse transcriptase (Takara), essentially as described previously (Suzuki et al. 1992). PCR was subsequently carried out with the sCT primers (Table 1) using a Program Temp Control System, PC-701 (Astec, Fukuoka, Japan). The amplification profile for 30 cycles was: dissociation at 94 °C for 1·5 min, annealing at 55 °C for 1·5 min and extension at 72 °C for 2·5 min. The final cycle included polymerization for 8 min. Amplification products were separated by electrophoresis and a major band was subcloned into pGEM-3Z vector (Promega, Madison, WI, USA). Sequencing reactions were conducted with a thermo sequenase cycle sequencing kit (USB, Cleveland, OH, USA), and nucleotide sequences were analysed using a Li-Cor automated DNA sequencer, model 4200L-2G (Li-Cor, Lincoln, NE, USA). A DNA probe was synthesized from a cDNA fragment encoding sCT-I, using a digoxigenin (DIG)-high prime kit (Roche Diagnostics GmbH, Mannheim, Germany).

DNA cloning and sequence analysis

Approximately 18 000 recombinants from the amplified cDNA library were screened by plaque hybridization. Hybridization with the above cDNA probe and post-hybridization washing were performed essentially according to the manufacturer’s instructions. Hybridization signals were detected with 25 mM CSPD, a 1,2-dioxetane chemiluminescent enzyme substrate (Tropix, Bedford, MA, USA).
USA), on Hyperfilm-ECL film (Amersham Biosciences, Piscataway, NJ, USA) after incubation with alkaline phosphatase-conjugated anti-DIG antibody (Roche). The pBK-CMV phagemid vectors with inserts were excised in vivo from the ZAP express vectors of positive recombinants, using the ExAssist helper phage (Stratagene). The nucleotide sequences of these DNAs were analysed using a Li-Cor automated DNA sequencer. A phylogenetic tree of CT precursors from the rainbow trout, chum salmon (Poeschl et al. 1987), puffer fish Fugu (Clark et al. 2002), chicken (Lasmoles et al. 1985), mouse (Rehli et al. 1996), rat (Jacobs et al. 1981) and human (Craig et al. 1982) was constructed on the basis of the amino acid sequence, using the Clustal W algorithm (Thompson et al. 1994).

### Tissue distribution of CT mRNAs

Total RNA was extracted from various tissues of rainbow trout, using TRIzol reagent (Roche). After treatment with deoxyribonuclease I (Takara), 10 µg total RNA were subjected to RT-PCR, as described above. Preparation of total RNA from rat tissues and its RT-PCR analysis were conducted similarly with the mCT primers (Table 1). As for the trout, a pair of specific primers were synthesized and used for detection of each of sCT-I, -IV and -V mRNAs (Table 1). The sCT-I primers were designed to recognize the sequences common to rainbow trout cDNAs for sCTs-Ia and -Ib (accession numbers AB094135 and AB094136 respectively; Hidaka et al. 2004), and the PCR amplification specific to each of sCT-I and -IV mRNAs was confirmed by restriction enzyme cleavage: PstI (Takara) for the former, and NspI (Roche) for the latter. Elongation factor 1-α (EF1-α) cDNA (BE669055) was also amplified with specific primers (Table 1), as an endogenous control. A 10 µl aliquot of each amplified product was electrophoresed through an ethidium bromide-stained 2% agarose gel and photographed with a FAS-III digital camera system. Semiquantitative analysis was carried out by calculating the relative amount of each sCT amplicon to the corresponding EF1-α band, using Scion Image software.

### Localization of sCT-I mRNA

In situ hybridization histochemistry was carried out on wax sections of the ultimobranchial glands. In situ hybridization of the tissue sections was performed as described previously (Suzuki et al. 1997). Antisense and sense RNA probes were prepared by in vitro transcription from the fragments of rainbow trout sCT-IV and -V cDNAs, using a DIG-RNA labelling kit (Roche).

### Results

#### Molecular cloning of sCT cDNA

There was a distinct cDNA band of approximately 100 bp observed after gel electrophoresis of RT-PCR products, which were amplified from the ultimobranchial gland mRNA using the sCT primers (Table 1). After isolation and subcloning of the cDNA fragment, the nucleotide sequences

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**Table 1 The nucleotide sequences of PCR primers**

<table>
<thead>
<tr>
<th>Primer</th>
<th>Sequence</th>
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<tbody>
<tr>
<td>sCT Sense</td>
<td>5′-CAAGCGTTGCTCAACCTCA-3′</td>
</tr>
<tr>
<td>sCT Antisense</td>
<td>5′-CGTTTCTTGGCAGGCTGCC-3′</td>
</tr>
<tr>
<td>rtCT-I Sense</td>
<td>5′-AATCTGACGGTCTCCCT-3′</td>
</tr>
<tr>
<td>rtCT-I Antisense</td>
<td>5′-TCAAAAAAGGGGATTAA-3′</td>
</tr>
<tr>
<td>rtCT-IV Sense</td>
<td>5′-ATTATAAGCTCTCTGTT-3′</td>
</tr>
<tr>
<td>rtCT-IV Antisense</td>
<td>5′-GAAGAAATTTATHTG-3′</td>
</tr>
<tr>
<td>rtCT-V Sense</td>
<td>5′-CTCTTTTTCACTTTAAATTTG-3′</td>
</tr>
<tr>
<td>rtCT-V Antisense</td>
<td>5′-GCTATTTTTAGATAAGTTTC-3′</td>
</tr>
<tr>
<td>mCT Sense</td>
<td>5′-TGAGGGCTCTAGCTTGCACA-3′</td>
</tr>
<tr>
<td>mCT Antisense</td>
<td>5′-TGTTGTTTGTCTCCAGTC-3′</td>
</tr>
<tr>
<td>sEF1-α Sense</td>
<td>5′-GGCTGGTTCAGGGATGGA-3′</td>
</tr>
<tr>
<td>sEF1-α Antisense</td>
<td>5′-ATTGGAGGGTCGTTCTTGCT-3′</td>
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coding for sCTs-I and -IV were determined by sequence analysis. The sCT-I-encoding fragment was then used as a probe to screen 18,000 recombinants from the rainbow trout cDNA library. About 1,080 positive CT clones were obtained from the rainbow trout, and sequence analysis revealed the complete nucleotide sequences of two distinct sCT cDNAs: one for sCT-IV and the other for a novel sCT, which we have designated as sCT-V (Figs 1 and 2), in addition to the previously isolated sCT-I cDNAs (AB094135, AB094136). No cDNA clones were obtained for sCTs-II or -III.

Rainbow trout sCT cDNA and precursor

The cDNAs for rainbow trout sCT-IV (AB094137) and -V (AB094138) precursors consisted of 925 and 915 bases respectively, excluding the poly(A) tail (Fig. 1). Both the precursors were predicted to encode a signal peptide of 25 amino acid residues, an N-terminal procalcitonin (proCT), a putative cleavage site Lys-Arg, calcitonin, a cleavage and amidation sequence Gly-Lys-Lys-Arg, and a C-terminal peptide of 18 amino acids (Fig. 2). The rainbow trout sCTs-IV and -V conserved a sequence for potential N-linked glycosylation (Asn88-Leu89-Ser90), although the biological significance of CT glycosylation has yet to be determined (Fig. 2). The sCT-V was identical to sCT-IV except for Met at position 8, and the sCT-V precursor exhibited a 90% identity with the sCT-IV precursor. A phylogenetic tree of CT precursors from the rainbow trout, chum salmon, puffer fish, chicken, mouse, rat and human was created using the Clustal W algorithm (Fig. 3).
Phylogenetic analysis of the CT precursors from various species, including rainbow trout, chum salmon, and chicken, reveals distinct clusters with the rainbow trout sCT-I precursor closely related to chicken CT precursors (Sasayama 1999, Suzuki et al. 1999b). The tree topology is consistent with the similarity among CT peptides from these species and further suggests that the CT genes of teleost and chicken might be more similar than estimated by phylogenetic analysis of CT regions (Clark et al. 2002).

**Figure 2** Comparison of amino acid sequences among the CT precursors of rainbow trout (accession numbers AB094135–AB094138), chum salmon (Y007665), puffer fish (Fugu; AJ309015), chicken (X03012), mouse (X97991), rat (V01228) and human (X00356). The sequences were aligned using Clustal W (Thompson et al. 1994). Gaps marked by hyphens (-) have been inserted to optimize homology. Conserved amino acids are shown in black boxes. The sCT-V is a novel peptide with Val of sCT-IV substituted by Met (arrow). The sCT-V precursor is most similar to the sCT-IV precursor, with 90% identity. The sCT precursors are 43–52% identical to mammalian CT precursors. Asterisks denote potential signals for proteolytic processing, and dots indicate a sequence for potential N-linked glycosylation.

Tissue distribution of sCT mRNAs

Calcitonin gene expression in rainbow trout tissues was first investigated by RT-PCR using the oligonucleotide primers that can amplify all the known sCT-encoding sequences. A cDNA band of the predicted length was observed in all the tissues examined, including the ultimobranchial gland (Fig. 4a). In contrast, RT-PCR analysis of CT mRNA in rat tissues indicated that rat CT gene expression was restricted to the thyroid containing CT-secreting C cells, brain, stomach, lung, adipose tissue, testis, ovary and kidney (Fig. 4b). Southern blot analysis using CT cDNA probes from the rainbow trout or rat confirmed the authenticity of those amplified bands (data not shown).

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Tissue distribution of sCT-I, -IV and -V mRNAs was further examined in the rainbow trout by RT-PCR using specific primers. The specificity of primers for each of the sCT cDNAs had been assessed by performing PCR with the sCT clones as templates. Although sCT-IV and -V cDNAs had sequences very similar to those of the rtCT-I
primers (Table 1), these primers amplified only sCT-I cDNA (Fig. 5a, lanes 1–3). It was verified that the amplified sCT-I cDNA fragment was cut with a restriction enzyme, PstI (Fig. 5a, lane 4), as predicted by sequence analysis. PCR with the rtCT-IV primers (Table 1) amplified sCT-V cDNA in addition to sCT-IV (Fig. 5b, lanes 2, 3). However, the PCR amplicon from sCT-IV cDNA was shown to be cut by another restriction enzyme, NspI (Fig. 5b, lane 4), indicating that the amplification of sCT-IV cDNA can be distinguished from that of sCT-V, and can be confirmed by this treatment. The rtCT-V primers (Table 1) amplified only the corresponding cDNA fragment (Fig. 5c), as they were synthesized on the basis of the sequence of insertion unique to sCT-V cDNA.

RT-PCR analysis for rainbow trout tissues using the rtCT-I primers, coupled with PstI digestion, revealed that sCT-I mRNA was expressed, not only in the ultimobranchial gland, but also in the testis, ovary and gills (Fig. 6). After NspI treatment of the PCR bands amplified with rtCT-IV primers, expression of sCT-IV mRNA was found in the testis in addition to the ultimobranchial gland (Fig. 6). A PCR band for sCT-IV mRNA was also detected in the pituitary, but the authenticity of this band could not be verified because the amounts of amplification products were insufficient for restriction digestion. Although the amino acid sequence of sCT-V peptide was nearly the same as that of sCT-IV (Fig. 2), the distribution of sCT-V mRNA was unique and was detected in most tissues examined, except the kidney, gills and blood cells (Fig. 6). In spite of the wide distribution of the sCT mRNAs, all were expressed in the greatest amounts in the ultimobranchial gland, as clearly shown by the quantification analysis (Fig. 6). RT-PCR examination further revealed that the sCT-I, -IV and -V mRNAs were all expressed in the ultimobranchial gland of a single trout (data not shown). The PCR amplicons from these sCT mRNAs were also observed in a rainbow trout of a different strain (data not shown), corroborating the existence of sCT-I, -IV and -V genes in the genome of this species.

Histological localization of sCTs-IV and -V mRNAs

The sites of sCT-IV and -V mRNA expressions were examined by in situ hybridization histochemistry with DIG-labelled antisense RNA probes. Hybridization signal for both sCT-IV (Fig. 7a) and sCT-V (Fig. 7c) mRNA was observed abundantly in the parenchymal cells of trout ultimobranchial gland. The sCT-IV mRNA was also detected at low levels in the Leydig cells of the testis (Fig. 7b).
Although RT-PCR analysis indicated the presence of sCT-V mRNA in various tissues (Fig. 6), we could not detect any distinct signals in their wax sections (data not shown), suggesting much lower expression levels of sCT-V mRNA in these tissues.

**Discussion**

In the present study, two distinct cDNAs encoding sCT-IV and a novel sCT-V were cloned from the ultimobranchial gland of rainbow trout. The initiation site for translation was predicted to be
Figure 6 Tissue distribution of sCT-I, -IV and -V mRNAs in the rainbow trout, determined by RT-PCR analysis. Total RNA (10 µg) from various trout tissues were reverse-transcribed and amplified by PCR with rtCT-I, -IV and -V primers (Table 1). Distinct bands for sCT-I mRNA were detected in the ultimobranchial gland, gonads and gills, whereas a prominent band for sCT-IV mRNA was seen for the testis and ultimobranchial gland. The authenticity of these bands was confirmed by specific digestion with restriction enzymes: PstI for sCT-I, and NspI for sCT-IV. In contrast, the expression of sCT-V mRNA was shown to occur in most tissues examined, but not in the blood cells. The relative levels of mRNA expression were normalized to EF1-α (upper panel), with the maximum value of each sCT defined as 10 000.
ATG at positions 1–3 in each cDNA (Fig. 1), because the sequence ACCATGG at positions −3 to 4 was identical with the consensus sequence for initiation sites in other eukaryotic genes (Kozak 1981). The deduced sCT-IV and -V precursors were composed of an N-terminal peptide, sCT and a C-terminal peptide, but the N-terminal peptide was predicted to be further cleaved into a signal peptide of 25 amino acids and an N-terminal proCT, by analogy with other teleost N-terminal peptides (Suzuki et al. 1998) and human preproCT (Russwurm et al. 1999). In view of the presence of sCT-I cDNAs (AB094135, AB094136), our sequence analysis suggests that three types of CT, i.e. sCTs-I, -IV and -V, could be produced in the rainbow trout. The RT-PCR analysis and in situ hybridization histochemistry further indicated strong expression of their mRNAs in the ultimobranchial gland, suggesting that sCTs-I, -IV and -V are all secreted into the circulation from this site. It was also shown by the RT-PCR that the sCT mRNAs were expressed in various tissues at lower levels. It has been reported that the mRNA encoding a possible CT receptor is present in a number of tissues from the flounder, such as the gills, brain, intestine, testis and ovary (Suzuki et al. 2000b). It has been also documented that CT receptor mRNAs are expressed in most mammalian tissues (Kuestner et al. 1994, Sexton et al. 1999). Therefore, receptors for sCT are likely to

Figure 7 In situ hybridization histochemistry for (a, b) sCT-IV and (c) sCT-V mRNAs in wax sections (6 µm) of trout tissues. (a) Parenchymal cells of the ultimobranchial gland showed an intense hybridization signal with the DIG-labelled antisense sCT-IV cRNA probe. (b) A weak signal for sCT-IV mRNA was also seen in the Leydig cells (arrows) of the testis. (c) A strong signal for sCT-V transcripts was evident throughout the ultimobranchial gland. (d) Control section for (c). Hybridization with the sense cRNA probe exhibited no signal on the ultimobranchial gland. Scale bar represents 50 µm.
exist also in those trout tissues in which sCT-I, -IV or -V mRNAs were detected, implying an autocrine or paracrine action (or both) of sCT.

As for mammals, there is evidence to suggest that CT has various physiological functions in addition to preventing hypercalcemia and encouraging bone mineralization (Sexton et al. 1999). For instance, Nakhla et al. (1989) indicated that CT could regulate testicular function because sCT stimulated the secretion of androgen-binding protein from primary Sertoli cell-enriched cultures and increased the concentrations of both androgen and estrogen receptors in TM4 cells that were derived from immature mouse Sertoli cells. In fish, CT-immunoreactive cells have been observed in the intestine of goldfish, and the population of these cells increased in response to a soup lacking calcium more than to calcium-enriched soup, although the plasma calcium concentration was highest in goldfish fed with the latter (Okuda et al. 1999). It is therefore proposed that CT cells in the intestine might affect the absorption of nutrients in this fish. With respect to salmonid fish, little is known about the local functions of sCT. However, our RT-PCR analysis has demonstrated the tissue distribution of sCT-I, -IV and V mRNAs in various trout tissues. Intriguingly, their patterns of tissue expression were remarkably different: sCT-IV mRNA was detected in the testis, whereas sCT-V mRNA occurred in most tissues examined. Expression of CT mRNA had already been reported in the gills of pink salmon and rainbow trout (Martial et al. 1994, Sasayama 1999), but our RT-PCR analysis revealed that the CT mRNA expressed in this tissue is only sCT-I. In addition, in situ hybridization histochemistry showed expression of sCT-IV gene in the Leydig cells of trout testis. The cells expressing sCT-V mRNA could not be localized in any tissues other than the ultimobranchial gland by this technique, but it has been the experience of many laboratories that in situ hybridization can be ineffectual in detecting the sites of expression of transcripts when their levels of expression are low (Nelhs et al. 1996, Jagger et al. 2000). Furthermore, an investigation using radio-immunoassay demonstrated lower concentrations of CT-like immunoreactivity in various tissues of the rainbow trout (Fouchereau-Peron et al. 1990), in some of which only sCT-V mRNA was detected by our RT-PCR analysis. Therefore, sCT-V mRNA seems to be expressed at very low levels in most tissues, as suggested by the RT-PCR. Taken together, our results suggest that sCTs-I, -IV and -V might be produced locally in a tissue-specific manner. Although there is no direct evidence, these sCTs might have different local functions, depending on their particular tissue distribution.

Much attention has been focused on proCT, because it is a specific marker for systemic inflammatory responses in humans (Russwurm et al. 1999, Whicher et al. 2001). It has also been reported that injection of proCT augments the mortality of hamsters suffering from sepsis (Nylen et al. 1998). It would be of great interest to examine whether pro-sCTs have similar effects in rainbow trout, even though the amino acid homology between fish and mammalian proCTs is less than that between their CTs.

Structural comparison with other CTs indicates that sCT-V is a new peptide in which the Val at position 8 of sCTs-I and -IV is replaced by Met. It is known that sCT-IV inhibits bone-slice resorption in a pit formation assay, and shows an even greater hypocalcemic potency than sCT-I in a rat bioassay (Schechroun et al. 1999). Because rat, rabbit and human CTs have Met at position 8, as occurs in sCT-V, sCT-V could prove to have some biological activities that might be used for the therapeutic treatment of bone disorders.

The present phylogenetic analysis of CT precursors demonstrates that rainbow trout sCT genes can be classified into two groups: one containing sCT-Ia and -Ib genes, and the other sCT-IV and -V genes. As rainbow trout is tetraploid (Ohno et al. 1968, Hiraoka et al. 1993), these two groups of sCT genes might have been generated by a genome duplication event in ancestral fish, and thereafter sCT-IV and -V genes might have diverged by gene duplication. In addition, it is possible that other sCT genes may exist in the genome of rainbow trout, because sCT-I, -IV and -V mRNAs were not detected in the kidney and blood cells in which sCT cDNAs were amplified by PCR using sCT primers corresponding to the conserved sequences flanking the sCT-coding region (Fig. 4a).

The RT-PCR analysis of the tissue distribution of sCT mRNAs showed that sCT-I, -IV and -V mRNAs were all expressed most abundantly in the ultimobranchial gland of rainbow trout. Nothing, however, is known about the molecular mechanisms that intensify sCT gene expression in this
site. Our elucidation of the sCT cDNAs has now made it possible to identify the entire sCT genes of the trout and determine whether homologous enhancer elements occur in the 5′-flanking DNA of these sCT genes. In mammals, luciferase reporter assays with HeLa and 44–2C cell lines revealed that an 18 bp element (5′-GGCAGCTGTGCAAA TCCT), located approximately 1 kbp upstream of the transcriptional initiation site of rat CT gene, has a strong enhancer activity (Tverberg & Russo 1993). A sequence nearly identical to this element is also present in the human CT/CGRP enhancer region (Broad et al. 1989, Peleg et al. 1990). These distal enhancers contain potential binding elements for helix-loop-helix (HLH) protein and octamer-binding protein designated OB2. It has been suggested that the major HLH complex can be a heterodimer of the ubiquitous upstream stimulatory factors-1 and -2 (Lanigan & Russo 1997). Although OB2 protein has not yet been cloned, mobility shift competition assays indicate the existence of this factor in the CT/CGRP-producing CA77 and 44–2C cells, but not in non-C cell lines such as HeLa, GH3, and Rat-1, suggesting that OB2 protein is a cell-specific activator. Therefore, OB2-like protein might exist in the ultimobranchial gland of rainbow trout, enhancing sCT gene expression.

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