Cloning and expression of a novel CREB mRNA splice variant in human testis

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Abstract

Identification of genes specifically expressed in adult and fetal testis is important in furthering our understanding of testis development and function. In this study, a novel human transcript, designated human testis cAMP-responsive element-binding protein (htCREB), was identified by hybridization of adult and fetal human testis cDNA probes with a human cDNA microarray containing 9216 clones. The htCREB transcript (GenBank Accession no. AY347527) was expressed at 2.35-fold higher levels in adult human testes than in fetal testes. Sequence and ntBLAST analyses against the human genome database indicated that htCREB was a novel splice variant of human CREB. RT-PCR-based tissue distribution experiments demonstrated that the htCREB transcript was highly expressed in adult human testis and in healthy sperm, but not in testes from patients with Sertoli cell-only syndrome. Taken together, these results suggest that the htCREB transcript is chiefly expressed in germ cells and is most likely involved in spermatogenesis.


Introduction

Spermatogenesis is a unique physiological process during which male germ cells undergo a series of differentiation steps leading to the production of mature haploid sperm cells. This process involves the coordinated expression of many genes with unique cellular and temporal specificities. One of the molecular mechanisms playing a major role in spermatogenesis is the cAMP-dependent signaling pathway, which involves modulation of a multigene family of transcription factors that contain the basic transactivation and DNA-binding domains (bZIP), such as cAMP-responsive element-binding protein (CREB), cAMP-responsive element modulator (CREM), and activating transcription factor-1 (ATF-1) (Habener et al. 1995, Scobey et al. 2001). CREB is a member of the bZIP family of transcription factors, which consists of two functionally distinct domains: a carboxyl-terminal dimerization and DNA-binding (bZIP) domain, and an amino-terminal transcriptional transactivation domain (Meyer and Habener, 1993, Andrisani 1999). Transactivation of gene transcription by CREB is dependent on the phosphorylation of a single serine within the phosphorylation domain (P box or kinase-inducible domain (KID)) by cAMP-dependent protein kinase A (PKA) (Gonzalez & Montminy 1989). CREB is expressed in nearly all tissues tested thus far. However, in the testis, investigators have identified a number of novel alternative exon splicings that result in the synthesis of mRNAs encoding multiple isoforms of CREB (Hoeflter et al. 1990, Ruppert et al. 1992, Waeger et al. 1993). In the rat testis, these alternatively spliced CREB mRNAs are spermatogenic, cycle-dependent and expressed during development of the germ and Sertoli cells (Daniel & Habener 1998), indicating that the CREB isoforms may be major players in spermatogenesis.

Previous cDNA microarray hybridization studies in our laboratory identified a number of known genes as being alternatively spliced in human testis, including reticulon, HsMCAK, DEAD-box protein and RAD23B; some of these spliceoforms were demonstrated to be spermatogenesis-related (Cheng et al. 2002, Yin et al. 2002, Zhou et al. 2002, Huang et al. 2004). In this paper, we report a novel spliceoform of CREB, designated htCREB (human testis cAMP-responsive element-binding protein), which was primarily expressed in human testis and was almost undetectable in testis samples from patients with Sertoli cell-only syndrome (there was a very weak band in patients 4 and 5). As compared with other spliceoforms of CREB, htCREB employs both alternative exons and
alternative splice site selection, with the resulting transcript lacking the basic transactivating region-associated Q2 domain. Taken together, these results suggest that htCREB is likely to inhibit transcription and may play an important role in human spermatogenesis.

Materials and Methods

Samples

Informed consent was received from either the participants or their kin, and the ethics committee of Nanjing Medical University (China) granted research approval prior to sample collection. Human testis from adult men (37 years old) and elderly (73 years old) men were obtained from the Body Donor Center (Nanjing Medical University), and fetal testes were obtained from accidentally aborted (as in road accidents) 6-month-old fetuses (Clinical Reproductive Center, Nanjing Medical University). Testis tissue samples from five individuals with Sertoli cell-only syndrome (SCOS) were acquired via biopsy, and healthy volunteers with proven fertility and normal semen quality (assessed by WHO criteria, 1999) donated ejaculated sperm. Macroscopic and histologic examinations showed there was every level of spermatogenic cells and Sertoli cells around seminiferous tubules in adult human testis and elderly testis; however, there were fewer spermatid and spermatozoa in elderly testis than in adult testis. Moreover, in fetal testis, only Sertoli cells and spermatogonia converged at the mediastinum.

Construction of human cDNA microarray

A total of 9216 positive phage clones were randomly picked from a human testis insert phage cDNA library (Clontech, H15503U) and PCR amplified. The resultant PCR products were spotted on nylon membranes to generate a human testis cDNA microarray. The cDNA microarray preparation and hybridization signal analyses were performed as previously described (Sha et al. 2002).

Array scanning and analyzing clones of interest

Fetal testis, adult testis and human spermatozoa probes were prepared by incorporation of 33P-labeled dATP in a reverse transcription reaction, using 2 µg purified mRNA as the template. The arrays were scanned, and the radioactive intensity of each spot was linearly scanned with a 65 536 gray-grade in a pixel of 50 µm and read out with array gauge software (Fuji Photo Film, Tokyo, Japan). After subtraction of the background from an area where no PCR product was spotted, clones with intensities over 10 were considered positive, and were picked, sequenced and analyzed as described (Sha et al. 2002, Wang et al. 2004). The generated sequences were subjected to BLAST analysis (www.ncbi.nlm.nih.gov), which identified one as htCREB, a novel human CREB mRNA splice variant in human testis (GenBank Accession no. AJ347527). Further GenBank sequence analyses were performed to determine the htCREB homologs and their chromosomal localization, and the nucleic and deduced amino-acid sequences of htCREB were analyzed with Gene Runner software (www.genruner.com) and SMART PROGRAM (Schultz et al. 1998).

Expression of the htCREB transcript in human testis and sperm

To determine expression of htCREB in various developmental stages of testis, htCREB-specific RT-PCR was carried out on cDNA from adult (age 37 years) testes, ejaculated sperm, old (age 73 years) testes and fetal testes, together with testes with SCOS. RNA was extracted from the various sources with Trizol reagent (Gibco BRL, Grand Island, NY, USA) and reverse-transcribed into cDNA with AMV reverse transcriptase (Promega). The various cDNAs were PCR amplified with htCREB-specific primers (P1, 5’ CCA GCC ATC AGT TAT TCA G3’ (forward, nt 262–280, located in exon C of htCREB); P2, 5’ AGA CTT CAG CAC TTC CTA C3’ (reverse, nt 519–537, located between exons F and H)) (Fig. 3) in 20 µl PCR reactions containing 10 x PCR buffer (2 µl), 25 mmol/l Mg2+ (1.5 µl), 2 mmol/l dNTPs (1.5 µl), Taq DNA polymerase (5 U/µl) (0.1 µl), distilled water (10.9 µl), 5 pmol primer (1 µl), and template cDNA (2 µl). The amplification conditions consisted of an initial denaturation at 94°C for 5 min, followed by 30 cycles of 94°C for 30 s, 52°C for 30 s and 72°C for 30 s, with a final extension at 72°C for 7 min. The PCR products were analyzed with 1.5% (w/v) agarose electrophoresis, and β-actin was amplified as the control. The PCR product of htCREB in adult testes was gel purified with a kit (QiAprep Spin Miniprep Kit; Qiagen) and cloned by the TA cloning system (Promega). The inserts were sequenced from one end by an automated DNA sequencer-ABI100 (model no. 377).

Tissue distribution of the htCREB transcript

For determination of htCREB tissue distribution, cDNAs from 16 different human tissues (heart, brain, placenta, lung, liver, skeletal muscle, kidney, pancreas, spleen, thymus, prostate, testis, ovary, small intestine, colon and leukocytes) from a commercial human multiple tissue cDNA panel (MTC Panels I and II, cat. nos. K1420-1 and K1421-1; Clontech) were amplified and analyzed as described above, with β-actin as the positive control.

Results

cDNA microarray hybridization

The htCREB hybridization intensities in the adult testes, fetal testes and spermatozoa samples were 52.36, 22.31 and 14.25 respectively (Fig. 1), indicating a 2.35-fold greater expression in the adult testes than in the fetal testes, with a low expression level in spermatozoa.
Analysis of the cDNA and putative protein sequences

Sequence analysis revealed that the htCREB transcript was 1650 bp long and consisted of eight exons. The deduced 230-amino acid htCREB protein had the conserved motifs characteristic of the bZIP family members (Fig. 2). BLAST searching of the human genome database (www.ncbi.nlm.nih.gov/genome/seq/HsBlast.html) localized the htCREB gene to human chromosome 2q34, and revealed that it showed high homology to two other full-length cDNAs (Accession nos. NM_004379, CREB-A; NM_134442, CREB-B) belonging to the CREB family. Further analysis revealed that all three were splice variants transcribed from the CREB gene at 2q34. Splicing comparison of htCREB with the other human CREB variants illustrated that exons B, C and E were identical in the three transcripts, while in htCREB, exon G was spliced out and exons A, F, H and I were shorter than in the other transcripts (Fig. 3). In addition, htCREB exons A and H employed alternative 3′ splice site selections, while exons F and I employed alternative 5′ splice site selections. SMART PROGRAM (Schultz et al. 1998) comparisons of the htCREB (AAQ24858) and other CREB (NP_004370, CREB-A, 327aa and NP_604391, CREB-B, 341aa) amino-acid sequences showed that htCREB lacked the Q2 domain found in the other two transcripts (Fig. 3).

Expression of htCREB transcripts in human testis and sperm

Our RT-PCR analyses revealed that htCREB transcripts were not detected in fetal testis, but were found in ejaculated sperm, and testes from adult and elderly men, though at relatively lower levels in the testis from elderly men (Fig. 4). Sequence results suggested that this band was the desired PCR product. Furthermore, htCREB was nearly undetectable in the testes of five SCOS patients except for patients 4 and 5 (Fig. 5).

Tissue distribution of htCREB transcripts

RT-PCR analysis of a multiple tissue cDNA panel revealed that the htCREB mRNA was highly expressed in human testis and weakly expressed in other tissues, such as lung and kidney (Fig. 6). Intriguingly, no expression signal was detected in human ovary, indicating that this alternative splice product is specific to the male.

Discussion

The CREB proteins are members of a multigene family of transcription factors involved in cAMP-mediated transcription regulation. The factors in this protein family contain the basic transactivation and basic-domain-leucine-zipper (bZIP) DNA-binding domains (Lou & Gagel 2001), with the transactivation domain divided into two regions: the phosphorylation box (P-box or KID) encoded by exons E and F, and two glutamine-rich Q1 and Q2 domains, which are chiefly encoded by exons C and G. Exons H and I make up the bZIP DNA-binding domains (Fig. 3). The CREB proteins induce target gene expression via their constitutive activation domains (CAD or Q2) and kinase-inducible activation domains (KID), which function synergistically in response to cellular signals (Goto et al. 2002). The KID stimulates transcription via a phospho(Ser133)-dependent interaction with the coactivator paralogs, CREB-binding protein and p300, whereas the Q2 domain recruits the TFIID complex via a direct association with hTAFII130 (Asahara et al. 2001, Chaudhary & Skinner 2001).

The CREB gene contains at least 12 exons, several of which are alternatively spliced to encode a variety of CREB isoforms. Exon D is an alternatively spliced exon of 42 bp encoding 14 amino acids. In this work, the other two CREB isoforms, CREB-A and CREB-B, used for comparison, were produced by exon D splicing in or out (Fig. 3). Both of these isoforms were widely expressed in many tissues (Berkowitz & Gilman 1990). In the testis, exons W (between G and H), Y (between D and E) and Z (between W and H) can be alternatively spliced, and all of them are testis-specific and are most strongly detected in germ cells. CREB is predominantly a positive modulator of the cAMP-responsive genes, but in the testis, alternative exon splicing additionally results in the expression of repressor CREB isoforms. When exon W is spliced into the CREB mRNA, termination of translation by stop codons within exon W permits in-frame translation to reinitiate at downstream initiation codons, resulting in the production of inhibitor CREB isoforms (I-CREBs). The I-CREBs compete
Figure 2 (Continued).
Figure 2 Nucleotide and deduced amino-acid sequence of htCREB. The single-letter code of the predicted amino-acid sequence is indicated below the nucleotide sequence from positions 44 to 736 (230 amino-acid residues). The start and termination codons are shaded. The tail signal is boxed, and the sequences used as the RT-PCR primers are underlined.

Figure 3 Comparison of amino-acid sequences between htCREB and two other human CREB isoforms: CREB-A and CREB-B. The coding regions, contained between the two arrows, are labeled and boxed, with colors showing corresponding exons and domains. The Q1 region of CREB-A is shorter than CREB-B due to exon D splicing out. htCREB lacks the Q2 region found in the other spliceoforms (green box). P1: upstream primer, P2: downstream primer.

Figure 4 (a) RT-PCR and electrophoresis indicate that htCREB is expressed in adult testis, ejaculated sperm and testis from elderly men, but not in fetal testis. (b) Amplification of β-actin as the positive loading control. Ma: marker; A: adult testis; S: ejaculated sperm; E: testis from elderly men; F: fetal testis.

Figure 5 RT-PCR of htCREB mRNA in patients with SCOS (1–5). (a) Expression of the htCREB transcript was nearly undetectable in patients 1–3 and weakly detected in patients 4 and 5. (b) Control amplification of β-actin in the corresponding patients. Ma: marker.
with CREB for binding to CREs (such as those located in the promoter of the CREB gene) and downregulate cAMP-stimulated gene expression. The I-CREBs are expressed at specific stages of spermatogenesis, predominately in spermatocytes, and may account for cell- and stage-specific repression of cAMP-regulated genes (Walker et al. 1996, 1998). Furthermore, insertion of human testis-specific exon Z after exon W abolishes the synthesis of one of the two inhibitor CREBs, due to the introduction of an in-frame stop codon within exon Z. The splicing in of exon Z may be part of a human-specific mechanism for regulation of cAMP-dependent regulatory pathways in spermatogenesis, by abolishing the expression of a CREB repressor (Girardet et al. 1996).

Here, we used a constructed human adult testis cDNA microarray to identify genes related to human testis development and spermatogenesis, and identified another novel alternative spliceoform of CREB (designated htCREB) in human testis. Hybridization of our human cDNA microarray with adult testis, fetal testis and sperm samples demonstrated that the expression level of the htCREB transcript differed in adult testis from fetal testis by more than twofold, and that the htCREB transcript was expressed in sperm (Fig. 1). Tissue distribution analyses indicated that htCREB was highly expressed in human adult testis but weakly expressed in the other tissues, such as lung and kidney (Fig. 6). Taken together, these results suggest that the htCREB transcript probably plays a role in human germ cell maturation, especially during spermatogenesis.

To further examine whether htCREB was chiefly expressed in germ cells, we first used RT-PCR to examine differential expression at various developmental stages: fetal testis, adult testis, elderly testis and human ejaculated sperm. The htCREB transcript was detected at relatively high levels in adult testis and human ejaculated sperm, and at lower levels in elderly testis, and was not detected in fetal testis (Fig. 4). This result appears to contradict the microarray data, which indicated a positive signal in the fetal testis experiment. As RT-PCR is a sensitive assay, we suggest that the positive microarray result could be due to hybridization of the probe with another, yet unknown, spliceoform of CREB that was not detected with the RT-PCR primers. Developmentally, spermatogenesis does not occur in male embryos. In adults, spermatogenic cells undergo successive mitotic, meiotic and postmeiotic phases, and then form mature sperm. In elderly men, spermatogenesis becomes less efficient, and the quantity and quality of sperm decrease. Thus, we suggest that in the human testis, htCREB may be primarily expressed in germ cells and may function in spermatogenesis. Accordingly, we explored htCREB expression in the testes of five SCOS patients and found that htCREB transcripts were nearly undetectable in such patients (Fig. 5). As SCOS is a condition of the testes in which only Sertoli cells occur in the seminiferous tubules, our results further suggest that htCREB, like the other identified spliceoforms of CREB, is primarily expressed in germ cells.

Next, we focused on examining the putative function(s) of this CREB spliceoform in spermatogenesis via structure and function analyses. Compared with the other spliceoforms CREB-A and CREB-B, htCREB employed both alternative exons and alternative splice site selection. Because exon G is spliced out, the deduced htCREB protein lacks the glutamine-rich Q2 region but contains Q1 and the other CREB domains (Fig. 3). CREB is a bifunctional transcription activator, exerting its effects through a constitutive activation domain (CAD or Q2) and a distinct KID (Felinski et al. 2001). Previous studies of CREB function have yielded conflicting results with regard to its activation of constitutive transcription. Deletion studies performed by Gonzalez et al. (1991) suggested that the NH2-terminal glutamine-rich (Q1) domain of CREB was important for basal activation of the somato­statin promoter in F9 cells. However, these studies did not establish that this domain was capable of providing basal activation alone, and subsequent results indicated that these results might be specific to the experimental model (Quinn 1993). Another study reported that contributions from both the Q1 and Q2 domains were important for basal activation, with Q2 being crucial for basal activity (Quinn 1993). Ferreri and coworkers subsequently showed that deletion of 10 amino acids from the Q2 domain
significantly reduced the interaction of CREB with hTAFII130, and they concluded that the CREB Q2 domain is fully competent to serve as a strong activator in vitro (Ferreri et al. 1994). Another CREB family member, CREM (cyclic-AMP responsive element modulator), was shown to produce PKA-responsive transcriptional activators, and also exists in isoforms that explicitly antagonize cAMP-dependent transcription (Foulkes et al. 1991, Laoide et al. 1993). Exon 9 of CREM (roughly equivalent to exon G of the CREB gene) also codes for a glutamine-rich domain. When exon 9 was inserted into the CREM mRNA by alternate splicing, the CREM protein switched from acting as a repressor to functioning as an activator during spermatogenesis (Foulkes & Sassone-Corsi 1992). In contrast, CREM proteins lacking Q2 may function as repressors, because they are unlikely to form productive interactions with proteins in the basal transcription complex (Ferreri et al. 1994). Thus, the functionality of hCREB may be considered analogous to that of CREM. With exon G spliced out, hCREB is likely to be a repressor, inhibiting transcription and downregulating cAMP-stimulated gene expression. During spermatogenesis, this novel I-CREB probably competes with the activator CREBs to determine the transcriptional rates of the vitally important cAMP-regulated genes (Walker et al. 1996).

In summary, the present study reports the identification of a novel spliceoform of CREB, hCREB, which is chiefly expressed in human male germ cells. With exon G spliced out, hCREB is likely to be the repressor responsible for downregulating CAMP-dependent gene expression. This identification and analysis of an additional testis-related CREB isoform improves our understanding of gene expression and regulation of spermatogenesis.

Acknowledgements
The work was supported by grants from National Project 973, China (no. G1999055901), and Chinese Natural Science Funds (no. 30170485).

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