



Cloning and Characterization of the cDNA Encoding the α -Subunit of cGMP-Phosphodiesterase in Canine Retinal Rod Photoreceptor Cells

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Rod photoreceptor cyclic GMP-phosphodiesterase (cGMP-PDE, EC 3.1.4.17) is a key enzyme in the phototransduction cascade of the vertebrate retina. The enzyme is composed of α - and beta-catalytic subunits, and two identical inhibitory gamma-subunits. Once the phototransduction cascade is initiated by the absorption of light by rhodopsin, activated cGMP-PDE rapidly hydrolyses cGMP, depletion of which shuts cGMP-gated cation channels in the plasma membrane. The consequent hyperpolarization of the photoreceptor outer segment represents a large signal amplification and generates the visual neural impulse. Aberrant function of cGMP-PDE is causally associated with retinal degenerative diseases in man and animals. Mutations in the genes for the α - and beta-subunits of cGMP-PDE (*PDEA* and *PDE6B*, respectively) cause retinitis pigmentosa (OMIM entry) in some human families [6,8,9]. Defects in *PDE6B* also cause retinal degeneration in the *rd* mouse [3,11] and the *rd1* dog [5,13,14]. Rod-cone dysplasia 2 (*rcd2*), which affects the collie dog, also represents a defect of retinal cyclic GMP metabolism since retinal cGMP levels are significantly elevated and cGMP-PDE activity is deficient [16]; however, *rcd2* is not caused by a defect in *PDE6B*[1]. Thus it is likely that the *rcd2* locus codes for either another cGMP-PDE structural subunit [*PDEA*, or *PDEG* (the gene for the gamma-subunit of cGMP-PDE)], or for one of the other proteins in the phototransduction cascade that activates cGMP-PDE. Since the canine chromosomal locations of the *rcd2* locus and the genes involved in phototransduction are not known, none of these candidate genes can be ruled out based on their map location relative to the *rcd2* locus.

In order to identify the mutation responsible for the *rcd2* disorder, we have begun to examine the different candidate genes that code for the phototransduction cascade proteins in the dog. The cDNAs for the α -subunit of cGMP-PDE (*PDE α*) have been cloned and characterized from man [12], mouse [2], and cow [10,12]. However, no information is available in the literature on the canine *PDE α* cDNA.

In this study we characterize the canine *PDE α* cDNA from normal dog, present evidence for usage of alternate polyadenylation sites to generate the two different transcripts described in multiple species [2,7,12], and compare the deduced amino acid sequences for conservation through evolution.

RESULTS

Initially two different segments of canine *PDE α* cDNA were obtained by reverse transcription (RT) and polymerase chain reaction (PCR) using total retinal RNA and consensus primer pairs (*PDEA-1/PDEA-2* and *PDEA-3/PDEA-4*; Table 1) based on the coding regions of known *PDE α* cDNA sequences from other species. These RT-PCR amplified fragments were cloned (*1-PDEA* and *2-PDEA*) in pCRII vector (Invitrogen; San Diego, CA) and sequenced to confirm authenticity of the retina-specific *PDE* cDNA sequence. From the confirmed canine

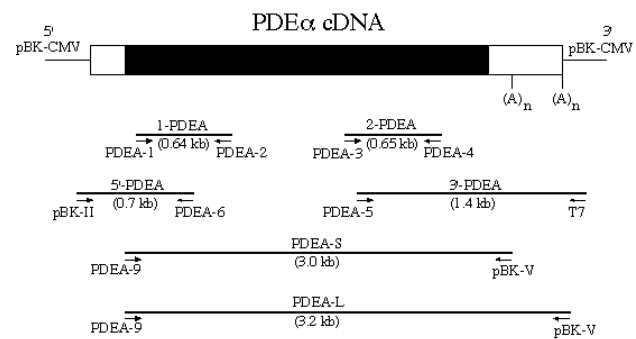


Figure 1. Strategy for cloning the canine *PDE α* cDNA. The cDNA containing the coding sequence (shaded box), the untranslated regions at the 5'- and 3'-end (open boxes) and two sites where poly (A) are added, is shown as an insert in the vector pBK-CMV (Stratagene, La Jolla, CA) used for the construction of the library. Clones containing different overlapping regions of the cDNA (*1-PDEA*, *2-PDEA*, *5'-PDEA*, *3'-PDEA*, *PDEA-S* and *PDEA-L*), the sizes of the cDNA fragments in the clones, and the primers (see Table 1 for sequence) used for generating those fragments are identified. The clones *1-PDEA* and *2-PDEA* were obtained by reverse transcription and polymerase chain reaction (RT-PCR) from canine retinal total RNA using the RT-PCR kit (Perkin-Elmer; Foster City, CA) as recommended by the manufacturer. Amplification of cDNA to obtain *1-PDEA* and *2-PDEA* was done for 30 cycles at 94 °C (1 min), 54 °C (2 min), 72 °C (2 min). Other clones (*5'-PDEA*, *3'-PDEA*, *PDEA-S* and *PDEA-L*) were obtained by screening a canine retinal cDNA library using a PCR based method. The PCR for screening the canine retinal cDNA library was done in 50 μ l containing 10 mM Tris-HCl, pH 8.3, 50 mM KCl, 0.2 mM dNTPs, 2.0 mM MgCl₂, 10% DMSO, and 1.25 units Taq polymerase (Life Technologies; Grand Island, NY). The conditions for PCR amplification of each fragment were as follows: *5'-PDEA* and *3'-PDEA* were obtained by 30 cycles at 94 °C (1 min), 60 °C (1.5 min), 72 °C (2 min); *PDEA-S* and *PDEA-L* were obtained by 30 cycles at 94 °C (1 min), 58 °C (2 min), 72 °C (3 min). All reactions were concluded with a single step extension reaction at 72 °C for 10 min.

PDE α cDNA sequence, new canine-specific primers were designed to amplify both the 5'- and 3'-ends of canine PDE α cDNA from a canine retinal cDNA library by PCR (Figure 1).

To amplify the 5'-end of the cDNA, forward vector-specific (pBK-II) and PDEA-specific reverse (PDEA-6) primers were used. From the PCR products, the largest fragment showing evidence for PDEA-specificity, based on PCR using internal primers, was cloned (5'-PDEA). To amplify the 3'-end of the cDNA, gene specific forward (PDEA-5) and vector specific reverse (T7) primers were used, and the amplified DNA fragment was cloned (3'-PDEA). The clone 3'-PDEA, however, lacked the poly (A) tail. We therefore designed a canine-specific primer (PDEA-9) from the 5'-end of the cDNA and used it in combination with the vector-specific reverse primer (pBK-V) for amplification of the entire open reading frame (ORF) and 3'-untranslated region (UTR) of PDE α cDNA from the cDNA library. The PCR resulted in amplification of two DNA fragments (3.0 and 3.2 kb), both of which hybridized to canine PDE α cDNA in Southern blots (data not shown). These two putative PDE α cDNA fragments (PDEA-L and PDEA-S) were cloned as described above.

Sequence of the PDE α cDNA was obtained from clones 1-PDEA, 2-PDEA, 5'-PDEA, 3'-PDEA, and PDEA-S, which contain overlapping fragments, from both directions. The identity of the larger clone (PDEA-L) as PDE α cDNA was confirmed by (a) partial sequencing (800 bp) of the 3'-end and two other upstream regions of the insert; (b) amplification of multiple overlapping fragments identical in size to those obtained from PDEA-S by PCR using the same set of primers; and (c) identical and predicted restriction enzyme digestion pattern of PCR amplified DNA fragments from both the clones (Figure 2). Sequences of all the primers used and their location in the canine PDE α cDNA or vector DNA are listed in

TABLE 1. SEQUENCE AND LOCATION OF PRIMERS USED FOR PCR

Primer sequence(5'to3')	Source of Primer	Name of primer	Location
gCTTTGCCAACAGTACTCAACC	hPDE α cDNA	PDEA1	192-215
AgTTTGTGcAGGTAACCTCAGGTG	hPDE α cDNA	PDEA2	815-836
GGCCCTGGTGGCGTTC	hPDE α cDNA	PDEA3	1759-1774
ATGGGATTTCTGTGCGACAC	hPDE α cDNA	PDEA4	2393-2412
TTCAACGTGGGGCAGACCAT	cPDE α cDNA	PDEA5	1832-1851
CACTACGTCCTTCCCATTCATTATGG	cPDE α cDNA	PDEA6	681-706
GTCATaAGAAGCTGTGCTTCTCC	cPDE α cDNA	PDEA7	380-404
GTCATGGGTGAGGTGACAGCAGAG	cPDE α cDNA	PDEA9	137-160
cCAACGTTTTGCCGAACCTCCAAG	cPDE α cDNA	PDEA11	2032-2054
TCCACCCTATTCTGGTCCCA	cPDE α cDNA	PDEA16	1036-1055
ATGGTCTGCCCCACGTTGAAGCC	cPDE α cDNA	PDEA17	1829-1851
TTGCTTGGCTGTCTGTGCTT	cPDE α cDNA	PDEA18	2624-2644
GCAGGTGCACACTAGTGGATCC	pBKCMV	pBKII	1092-1113
CCGCTCTAGAAGTACTCTCGAGTT	pBKCMV	pBKV	1052-1067
CGACTCACTATAGGGCAATT	pBKCMV	T7	980-1000

Primers (PDEA-1 to PDEA-4) used for RT-PCR correspond to the human (h) PDE α cDNA sequence, selected from the consensus region in different species. All other PDEA primers correspond to canine (c) PDE α cDNA sequence. Location of all the PDEA primers are shown with respect to cPDE α cDNA sequence. Nucleotides (shown in lower case) in some primers have mismatches with the canine sequence because those primers were selected either from the human sequence, or from preliminary canine sequence. Phagemid vector (pBK-CMV) specific primers were selected either from the multiple cloning site or from the flanking region. The bold-italicized region of primer pBK-V represents the linker used to make the canine cDNA library.

Table 1.

The two full length canine rod PDE α cDNAs are 2988 and 3233 nucleotides long, including the poly (A) tail (Figure 3). We have previously reported the presence of two distinct PDE α mRNAs, 3.3 and 3.0 kb, expressed in equivalent amounts in canine retina by northern blot [7]. Thus the sizes of the canine PDE α cDNAs described here (GenBank accession number U52868) are in complete agreement with our observation from northern analysis [7]. We have not formally determined the transcription start site; however, the clone containing the most upstream 5'-noncoding region contains 139 nucleotides in the 5'-UTR followed by the ATG codon for initiation of translation. The stop codon (TAA) corresponds to positions 2723 through 2725 of the cDNA sequence. The 3'-UTRs of the 3.0 kb and 3.3 kb transcripts, represented by clones PDEA-S and PDEA-L, are 263 and 508 nucleotides respectively. A non-consensus putative polyadenylation signal (ATTAAA) is present in the corresponding locations of the smaller (PDEA-S) and larger (PDEA-L) clones. In PDEA-S this first polyadenylation signal (Figure 3, nucleotides 2948 through 2953) is 15 nucleotides upstream of the poly (A) tail.

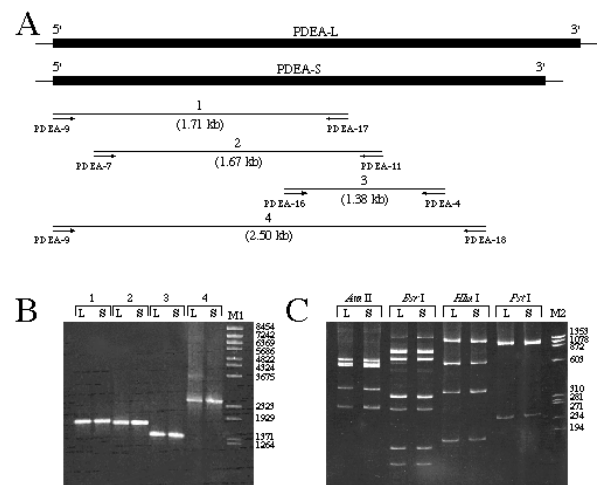


Figure 2. Identity of PDEA-L clone as PDEa cDNA sequence. (A) Relative size of the inserts in the PDEA-L and PDEA-S clones, the DNA fragments amplified by PCR, primers (see Table 1 for sequence) used, and the predicted sizes of the amplified fragments are shown. PCR condition for amplified fragments are as follows: fragments 1 and 2 were obtained by 30 cycles at 94 °C (1 min), 64 °C (1 min), 72 °C (2 min); fragment 3 was obtained by 30 cycles at 94 °C (1 min), 54 °C (1 min), 72 °C (1.5 min); and fragment 4 was obtained by 35 cycles at 94 °C (1 min), 58 °C (2 min), 72 °C (3.5 min). All reactions were concluded with a single step extension reaction at 72 °C for 5 min.

(B) Agarose gel (0.8%) electrophoresis of PCR products from PDEA-S (S) and PDEA-L (L) clones. Numbers 1 through 4 identify DNA fragments amplified from regions of the clones represented in panel A. Lane M1 represents BstE II-digested lambda DNA markers (C) Polyacrylamide gel (6%) electrophoresis of restriction enzyme digestion of PCR product (4 in panel A) from PDEA-S (S) and PDEA-L (L) clones. Lane M2 represents Hae III-digested phi X 174 markers. The length of the DNA fragments (bp) in both markers (M1 and M2) are shown next to the marker lanes. The PCR and the restriction enzyme digestion results indicate that both the clones are identical in the regions examined.

sites. The same mechanism could well account for the observed different transcript sizes within other species. The canine sequence does not contain the canonical polyadenylation signal (AATAAA); instead a similar sequence motif (ATTAAG) was identified in both appropriate locations (Figure 3). This alternative motif has been demonstrated to serve as a surrogate polyadenylation signal with ~80% efficiency [15]. A non-consensus polyadenylation signal (AATACA) has also been reported to be present in human PDE α cDNA [12].

It is noteworthy that while PDE α cDNA clones from 3 other species contain an ORF capable of coding for a polypeptide of 859 amino acids, the canine PDE α polypeptide is predicted to contain 861 amino acids. With respect to the human and mouse sequences, the two extra amino acids are located as the 849th (Ala) and 850th (Gly) residues (Figure 3). Similar to the comparison made between human and bovine PDE α sequences [12], we noted that the differences in amino acid sequence among the four species (human, mouse, bovine, and canine) are clustered in the 225 N-terminal and 45 C-terminal residues (data not shown). The conserved region includes the domain present in several eukaryotic cyclic nucleotide phosphodiesterases [4]. Alignment of amino acid sequences of canine PDE α and PDE β shows a 72% overall identity, and the domains that are most dissimilar are at the N-terminus (first 50 residues) and C-terminus (last 30 residues).

The data presented here have been used to identify the possible mechanism for the presence of two transcripts for PDE α in different species, and to compare nucleotide/amino acid identity among rod-specific PDE α sequences of these species. Characterization of the wild type canine PDE α cDNA sequence will allow us to detect PDEA mutations in dogs affected with *rcd2* or other inherited retinal degenerations. Because collie dogs affected with *rcd2* have elevated retinal cGMP levels secondary to low PDE activity, we are currently investigating the possibility of a PDE α mutation in this disease.

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The typographical corrections below were made to the article on the date noted. These changes have been incorporated in the article and the details are documented here. 29 March 1999: The text "retinitis pigmentosa (RP)" was changed to "retinitis pigmentosa (OMIM entry 268000)" in the first paragraph of the introduction. The abbreviations "Fig." and "no." were expanded to "Figure" and "number", respectively, throughout the article. The characters enclosing citations to references were changed from parentheses () to brackets []. Several instances of incorrect spacing between a word and punctuation have been corrected.