Identification and characterization of sea squirt telomerase reverse transcriptase

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Abstract
Telomerase is essential for maintaining telomere length and chromosome stability in most eukaryotic organisms. The telomerase ribonucleoprotein complex consists of two essential components, the catalytic telomerase reverse transcriptase protein (TERT) and the intrinsic telomerase RNA. The sea squirts, as urochordates, occupy a key position in the phylogenetic tree of the chordates: they diverged from the other chordates just before the lineage of vertebrates, and thus provide special insight into the origin and evolution of vertebrate genes. Here, we report the cloning and characterization of TERT genes from two sea squirts, Ciona intestinalis and Ciona savignyi. The C. intestinalis TERT (Cin TERT) gene encodes 907 amino acids and consists of 17 exons, which are similar to vertebrate TERT genes. The C. savignyi TERT (Csa TERT) gene encodes 843 amino acids, but surprisingly does not contain any introns. Both Ciona TERTs contain all of the reverse transcriptase (RT) motifs (1, 2, A, B, C, D, and E) that are typically present in telomerase and viral RTs. Interestingly, the alignment of Ciona and vertebrate TERT sequences reveals a previously unknown motif, named motif 3, located between motifs 2 and A. The Ciona TERT gene is expressed in all tissues analyzed except the brain and heart. This is the first report of the TERT gene in invertebrate chordates.

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1. Introduction
Telomeres are DNA–protein complexes that cap the ends of linear chromosomes to prevent end-to-end fusion or apoptosis (Blackburn, 2001; deLange, 2002). The telomeric DNA in vertebrates consists of tandem repeats of TTAGGG sequence, and associates with a number of telomere-binding proteins. In addition to their protective role in chromosome stability and genomic integrity, telomeres also play a critical role in cellular aging. In normal somatic cells, telomeres shorten progressively, which eventually leads to cell senescence or death. Immortal cells, such as stem cells and tumor cells, utilize an enzyme called telomerase to maintain telomere length and cellular immortality.

Telomerase is a specialized reverse transcriptase (RT) that adds telomeric DNA repeats to chromosome ends (Greider and Blackburn, 1985, 1987). The telomerase enzyme consists of two essential components, the catalytic telomerase reverse transcriptase (TERT) protein, and the telomerase RNA (TR) that provides a template for telomeric DNA synthesis (Greider and Blackburn, 1989). The TERT protein was cloned initially from the ciliate Euplotes aediculatus and the yeast Saccharomyces cerevisiae (Lingner et al., 1997). Homologs of TERT have since been identified in a variety of eukaryotic species, including vertebrates — human, dog, mouse, rat, hamster, chicken, Xenopus and Fugu fish (Meyerson et al., 1997; Nakamura et al., 1997; Greenberg et al., 1998; Guo et al., 2001; Kuramoto et al., 2001; Wong et al., 2003; Delany and Daniels, 2004; Nasir et al., 2004; Yap et al., 2005); plants — rice, Asparagales and Arabidopsis (Fitzgerald et al., 1999; Oguchi et al., 1999; Heller-Uzsynska et al., 2002; Sykorova et al., 2006); yeasts — Schizosaccharomyces

Abbreviations: aa, amino acids; bp, base pairs; cDNA, complementary DNA; TERT, telomerase reverse transcriptase; kb, kilobase(s); kDa, kilodalton (s); PCR, polymerase chain reaction; RACE, rapid amplification of cDNA ends; RT, reverse transcriptase; RT-PCR, reverse transcriptase-polymerase chain reaction; TR, telomerase RNA; UTR, untranslated region.

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**2. Materials and methods**

**2.1. Total RNA isolation**

Total RNA was isolated from various tissues (brain, heart, intestine, stomach, ovary, muscle and testis) of *Ciona intestinalis* using Trizol reagent (Life Technologies, USA). Live animals of *Ciona intestinalis* were obtained from Marine Biological Laboratories (Woods Hole, Massachusetts, USA). In brief, 30–50 mg of fresh tissue from *C. intestinalis* was homogenized in 1 ml Trizol reagent and the resulting solution was mixed with 0.2 ml chloroform, followed by centrifugation at 12,000 × g for 15 min at 4 °C. The aqueous phase containing total RNA was transferred to a new tube and extracted once or twice with acid-phenol/chloroform, pH 4.5 (Ambion, USA), followed by chloroform extraction and isopropanol precipitation. RNA concentrations were determined by optical density (OD$_{260}$) measurement using a Nanodrop ND-1000 spectrophotometer (NanoDrop Technologies, USA).

**2.2. Identification and cloning of *Ciona* TERT genes**

The *C. intestinalis* genome database (Assembly version 2.0 at a coverage of 11×) at http://genome.jgi-psf.org was searched using the TBLASTN algorithm with the vertebrate TERT protein sequences as query. A scaffold (scaffold #497) containing putative exons that are homologous to vertebrate TERT genes was identified. Gene-specific primers were designed to PCR amplify partial cDNA fragments of *Ciona* TERT. The exact 5′ and 3′-ends of the full-length cDNA sequence of the *C. intestinalis* TERT gene (named CinTERT) were determined by 5′- and 3′-rapid amplification of cDNA ends (RACE) using a SMART RACE cDNA Amplification Kit (Clontech, USA) following manufacturer’s instructions. To confirm the cDNA sequence, primers annealed to the 5′ and 3′-UTR regions were used for RT-PCR, and the DNA product of RT-PCR was sequenced directly. The exon and intron boundaries of the CinTERT were determined by aligning the cDNA sequence to the genomic sequence of the scaffold #497. The cloned CinTERT cDNA sequence has been deposited in the GenBank database under the Accession No. EF707623.

The *Ciona savignyi* genome database (13× coverage) at http://www.broad.mit.edu/annotation/Ciona was searched using the CinTERT sequence as query. A single ORF that encodes a putative TERT sequence (843 amino acids) was found in the scaffold #427. The *C. savignyi* genomic DNA and cDNA samples were obtained from Dr. William Smith at U.C. Santa Barbara. The CsATERT gene was PCR amplified from genomic DNA and cDNA samples using a pair of primers, a forward primer (5′-ACG ACC TGA ATA TCT TGG ATT G3′) and a reverse primer (5′-ACC TTC TCA ACC TTG GAT G3′) and 3′-UTR regions were used for RT-PCR, and the DNA product of RT-PCR was sequenced directly. The exon and intron boundaries of the CinTERT were determined by aligning the cDNA sequence to the genomic sequence of the scaffold #497. The cloned CinTERT cDNA sequence has been deposited in the GenBank database under the Accession No. EF514225.

**2.3. Detection of *Ciona* TERT expression by RT-PCR**

Approximately 1 μg of total RNA was reverse transcribed into complementary DNA (cDNA) using Thermoscript (Invitrogen, USA) and an oligo dT18 primer. The cDNA was then used as template in PCR reactions with Ex-Taq DNA polymerase (Takara Bio Inc. Japan) and specific primers. A pair of primers complementary to CinTERT exon 15 (5′-CAA CCA TGG GTG ATT GGG AAT CCA-3′) and exon 17 (5′-GGG ACA TTC CAA CTG CCC TTT GTA G-3′) was used to amplify the TERT cDNA from exons 15 to 17. Another pair of primers complementary to exon 5 (5′-ACC TTC TCA ACC TGG ATT G3′) and 3′-UTR regions were used for RT-PCR, and the DNA product of RT-PCR was sequenced directly. The exon and intron boundaries of the CinTERT were determined by aligning the cDNA sequence to the genomic sequence of the scaffold #497. The cloned CinTERT cDNA sequence has been deposited in the GenBank database under the Accession No. EF514225.
primers, a forward primer (5′-AGA GCT ACG AAC TTC CTG ACG GAC AG-3′) and a reverse primer (5′-GAA CAT AGT GGA ACC TCC AGA GAG AAC-3′), complementary to exon 4 and exon 5, respectively. The PCR was carried out with 95 °C for 2 min, followed by 30–40 cycles of 95 °C for 15 s, 55 °C for 30 s, and 72 °C for 1–2 min, followed by a final elongation at 72 °C for 3 min. The more abundant actin cDNA was amplified with only 30 cycles of PCR, whereas the TERT cDNA was amplified with 40 or 45 cycles of PCR.

2.4. Sequence analysis of Ciona TERT

Multiple sequence alignment of TERT proteins was performed using ProbCons for the RT domain and C-terminal region. ProbCons is an improved tool for multiple sequence alignment of proteins using a combination of probabilistic modeling and consistency-based alignment techniques to achieve a higher accuracy (Do et al., 2005). ClustalX was used to align the N-terminal region in which the sequence and length vary significantly (Chenna et al., 2003). A phylogenetic tree was constructed from the aligned sequences of the RT domain using MEGA version 3.1 obtained from http://www.megasoftware.net (Kumar et al., 2004).

3. Results and discussion

3.1. Identification and characterization of the sea squirt TERT (Ciona TERT) genes

To find the TERT gene in the C. intestinalis genome database, we performed a TBLASTN search using the vertebrate TERT protein sequences as query. The search successfully identified a putative Ciona TERT sequence that encodes a conserved RT motif B (GIPQG). Gene prediction analysis of this sequence as well as its flanking sequences obtained from the genome database exhibited a number of small and scattered open reading frames (ORFs) separated presumably by introns. To obtain the full-length cDNA sequence of this gene, we carried out 5′- and 3′-RACE using a cDNA library prepared from C. intestinalis ovary mRNA and gene-specific primers that target the predicted exon sequences of the putative Ciona TERT gene.

The 5′- and 3′-RACE revealed a cDNA sequence of 3588 bp that contains a 5′-UTR of 402 bp, a 3′-UTR of 462 bp and an ORF of 2724 bp assuming that translation initiates at the first ATG codon (position 403–405). This ORF is predicted to encode a protein of 907 amino acids. On the basis of sequence similarity with other TERT proteins, we have named this protein gene the C. intestinalis TERT (CinTERT). To characterize the promoter region of the TERT gene, we performed PCR to amplify the upstream flanking region of the gene. About 500 bp of the CinTERT promoter sequence was obtained. A putative TATA box, 5′-TATTAA-3′, is located 36 bp upstream of the transcription initiation site (Fig. 1B, underlined). Based on the result of the 3′-RACE, polyadenylation of the CinTERT mRNA occurs at the A triplet. A putative polyadenylation signal, 5′-AAAAATAA-3′, is located 20 bp upstream of the A triplet (Fig. 1C, underlined).

To predict more precisely the translation initiation site, the cDNA sequence was analyzed using a recently developed algorithm that predicts translation initiation sites with high sensitivity and specificity (Li and Leong, 2005). Using this algorithm, the same ATG codon at position 403–405 was predicted with the highest probability to be the translation initiation site (data not shown). Mapping of the cloned cDNA sequence to the genomic sequence from the database indicated that the CinTERT is composed of 17 exons and 16 introns (Fig. 1A), similar to vertebrate TERT genes (Leem et al., 2002; Yap et al., 2005). All introns conformed to the GT/AG rule at the splicing junctions and have sizes ranging from 239 to 897 bp, similar to the sizes of introns in vertebrate TERT genes. From the transcription start site to the polyadenylation site, the genomic sequence of the CinTERT gene spans 10.5 kb, smaller than the 41 kb human TERT gene (Leem et al., 2002) but larger than the 6.3 kb fugu gene (Yap et al., 2005). The CinTERT gene was mapped to chromosome 8q based on a BLAST search of the C. intestinalis genomic and cDNA database (http://ghost.zool.kyoto-u.ac.jp) (Satou et al., 2005). Overall, the similarity in protein sequence and gene structure suggests that the CinTERT identified here is an ortholog of vertebrate TERTs.

Two types of C. intestinalis have recently been reported (Suzuki et al., 2005; Nydam and Harrison, 2007). One type inhabits the Pacific Ocean, while the other type inhabits the northwest Atlantic Ocean. The CinTERT cDNA cloned in this study was from an Atlantic C. intestinalis (see Materials and

Fig. 1. Schematic of C. intestinalis TERT gene structure. (A) Splicing pattern of C. intestinalis TERT gene. Exons are numbered from 1 to 17, and shown as open boxes, whereas introns are shown as lines. The 5′ and 3′-untranslated regions (UTRs) are shown. The sizes (bp) of each intron and exon are shown. (B) Transcriptional start site sequence of Ciona TERT gene. The transcription start site is shown in bold letters and indicated with +1. The putative TATA box, 36 bp upstream of the start site, is underlined. (C) Polyadenylation site sequence of Ciona TERT gene. Polyadenylation sites are shown in bold letters with the cleavage sites indicated by arrowheads. A putative polyadenylation signal is underlined.
methods), while the whole-genome sequences were derived from a single Pacific *C. intestinalis* captured in Half Moon Bay, California, (Dehal et al., 2002). Between these two *Cin* TERT sequences, we observed an 89% identity at the nucleic acid sequence level and 87% identify (or 91% homology) at the protein sequence level. A recent report showed that the two types of *C. intestinalis* differ by 12% in the sequences of a mitochondrial gene, cytochrome c oxidase subunit I (Nydam and Harrison, 2007). It has also been observed that the *Ciona* genome is highly polymorphic (up to 1.2% of heterozygosity) (Dehal et al., 2002). To examine the individual polymorphism, we performed direct sequencing of RT-PCR products amplified independently from three different Atlantic *C. intestinalis* individuals. The cDNA sequences obtained from all three individuals were nearly identical (data not shown). The *Cin*-TERT gene discussed below refers to the one identified experimentally from Atlantic *C. intestinalis* in this study.

In addition to the *C. intestinalis*, *C. Savignyi*, a close relative of *C. intestinalis*, also has had its genome sequenced (Vinson et al., 2005). Using the identified *Cin* TERT protein sequence as a query, we searched the *C. savignyi* genome database and a single ORF that encodes a putative TERT sequence was found. Surprisingly, the deduced sequence of *C. savignyi* TERT (named *Csa* TERT) encodes an 843 amino acid protein that contains all conserved motifs found in *Cin* TERT and vertebrate TERTs. Finding a near full-length ORF in the genomic sequence suggests that this putative *Csa* TERT gene contains no introns. The absence of introns in the *Csa* TERT gene was confirmed by sequencing PCR products amplified from both genomic DNA and cDNA (data not shown). Based on a pairwise sequence alignment, the 5′-end of the *Csa* TERT ORF was mapped to exon 2 of the *Cin* TERT gene. The *Csa* TERT protein is, thus, 36 amino acids shorter than the *Cin* TERT protein at the N-terminal region (Fig. 2A). The entire *Cin* TERT and *Csa* TERT protein sequences share 47.4% identity and 63.0% similarity. Because the *Csa* TERT possesses all motifs essential for telomerase activity and lacks features typical for a pseudogene such as a 3′-terminal poly(dA) tail, this intron-less *Csa* TERT gene is not likely to be a pseudogene. More rigorous BLAST searches of the *C. savignyi* genome did not reveal any other candidate sequences for the TERT gene. Although we speculate that the *Csa* TERT gene might have resulted from Fig. 2. (A) Multiple amino acid sequence alignment of the N-terminal region of TERT proteins. The alignment includes TERT sequences from human (*Homo sapiens* AF015950), dog (*Canis familiaris* AF80351), mouse (*Mus musculus* NM-009354), hamster (*Mesocricetus auratus* AF149012), *Xenopus* (*Xenopus laevis* AF212299), fugu fish (*Fugu rubripes* AY861384), sea urchin (*Strongylocentrotus purpuratus* EF611988), sea squirts (*C. intestinalis* and *C. savignyi*) and nematodes (*Caenorhabditis elegans* NP-492374, *C. remanei* DQ178631 and *C. briggsae* CAE66970). Shaded residues represent the conserved amino acids. The positions of amino acids are shown at both ends of each line. Hyphens indicate gaps introduced to optimize alignment. Identical residues are shown above the alignment. Chicken (*Gallus gallus* AY502592) TERT sequence is omitted from this alignment due to its unusual length variation in the N-terminal region. (B) Multiple amino acid sequence alignment of the central and C-terminal regions. Conserved residues in the T motif and RT motifs are shaded with specific colors and indicated above the alignment. Residues absolutely conserved in all RTs are indicated with red asterisks below the alignment. (For interpretation of the references to colour in this figure legend, the reader is referred to the web version of this article.)
Fig. 2 (continued).
gene duplication or RNA-mediated retrotransposition, the mechanistic basis for the origin of this intron-less TERT gene remains to be elucidated.

3.2. Comparison of Ciona and vertebrate TERT protein sequences

Because of the different degrees of sequence and length conservation in different domains of the TERT protein, a given sequence alignment algorithm sometimes fails to align the protein sequences in different regions with the same accuracy. Different domains of the TERT protein might have evolved at different rates along different lineages of species. For example, the sequences and length of the N-terminal and C-terminal domains are usually more variable and are thus more difficult to be aligned correctly. To achieve a more accurate sequence alignment to reveal important motifs, we employed a novel approach with which sequences of individual domains of the protein from closely related species were aligned separately.

The sequences of the N-terminal domain of TERTs from vertebrates, sea squirts, sea urchin and nematodes were aligned using the ClustalX program. The N-terminal domain of vertebrate TERT proteins contains five telomerase-specific motifs, named region v-I, v-II, v-III, v-IV and motif T (Kuramoto et al., 2001). Similar to the Ciona TERT, nematode TERTs also have a truncated N-terminal domain (Malik et al., 2000; Meier et al., 2006). By aligning the sequences of the N-terminal domain from sea squirts, nematodes and sea urchin TERTs, we found that both squid and nematode TERTs lack region v-I but contain region v-II near the N-termini of the proteins (Fig. 2A).

Region v-I has been previously defined as motif GQ (Xia et al., 2000; Bosoy et al., 2003), and is conserved in most species with the exception of insects (Osanai et al., 2006) and nematodes (Malik et al., 2000). The GQ motif has been shown to be important for repeat addition processivity in human and yeast TERTs (Moriarty et al., 2004; Lue, 2005). It also shows a strong affinity to the single-stranded telomeric DNA primer (Jacobs et al., 2006). Within the region v-I-GQ, there is another motif called the “dissociate activities of telomerase” (DAT) motif that has been shown to be required for telomerase activity in vivo (Armbuster et al., 2001). It would be interesting to know how Ciona telomerase functions without these important motifs in the region v-I.

While lacking region v-I, the Ciona TERT contains most residues conserved in region v-II, v-III and v-IV (Fig. 2A). Region v-III includes a previously identified motif CP that is conserved in ciliates (Bryan et al., 1998). Region v-IV has been previously defined as motif QFP, and is also conserved in Ciona TERT (Fig. 2A). The nematode TERTs are much more divergent in sequence and length, and contain only parts of the region v-III, v-IV and motif T (data not shown) (Meier et al., 2006).

The sequence alignment of chordate TERTs, in addition to identifying chordate-specific motifs, also refines the boundaries of known RT motifs of TERT proteins. The universal RT motifs (1, 2, A, B, C, D and E) were originally identified based on RT sequences in viruses or retrotransposons (Poch et al., 1989; Xiong and Eickbush, 1990), and later on TERT sequences (Lingner et al., 1997; Nakamura et al., 1997). However, the boundaries of these motifs in TERT are not well defined and can vary between different groups of species. With the appropriate evolutionary distance between Ciona and human, the alignment of chordate TERT sequences allows for a better boundary definition of the conserved motifs in human TERT. In this study, the motif boundaries are defined by the first and last conserved residues within the motif (Fig. 2).

To obtain a more accurate alignment of the sequences from the RT and C-terminal domains, we employed a novel sequence alignment algorithm, ProbCons, that is based on probabilistic consistency and has been shown to have significant improvement over other methods for multiple sequence alignment of proteins (Do et al., 2005). The central domain of TERT contains all previously known RT motifs (i.e. 1, 2, A, B, C, D and E) (Fig. 2B). In Ciona TERTs, the distance between motifs A and B is significantly shorter than other TERTs. Ciona TERTs contain all invariant amino acids that are conserved in RTs, including a lysine (K) in motif 1, an arginine (R) in motif 2, an aspartic acid (D) in motif A, a glutamine (Q) and a glycine (G) in motif B, two aspartic acids (DD) in motif C, a lysine (K) in motif D and a glycine (G) in motif E (Fig. 2B). These invariant residues are crucial for enzyme activity (Lingner et al., 1997). In addition to the previously known RT motifs, we have also identified a novel motif, named motif 3, located between motifs 2 and A (Fig. 2B). This motif 3 is also present in other groups of species outside of chordates such as sea urchin, plants and ciliates (data not shown). The sequences of motif 3 from human and Ciona share 28.6% identity and 45.7% similarity in sequence. This sequence conservation suggests that motif 3 plays an important role in telomerase function. Mutations of the conserved residues in the motif 3 altered activity or processivity of human telomerase (Xie and Chen, unpublished data).

Fig. 3. Schematic representation of TERT protein motifs. The structures of human (Homo sapiens), fugu (Fugu rubripes) and Sea squirt (Ciona intestinalis) TERT proteins are shown. Motifs are shown as boxes. Three domains, the N-terminal, the central-RT and the C-terminal domains are indicated above the schematic of the motif structures. The length (a.a), predicted molecular weight (MW, in KDa), and isoelectric point (pl) are indicated at the right.
The sequence alignment of the C-terminal region revealed a number of insertions and deletions between conserved motifs in the *Ciona* TERT (Fig. 2B). These gaps in the alignment allow better definition of the boundaries of each conserved region. We have thus redefined three conserved chordate motifs, named region c-1, c-2, and c-3, in the C-terminal region (Fig. 2B). These three regions contain most of the conserved residues that were previously shown to be critical for human telomerase activity *in vivo* (Banik et al., 2002).

Overall, motifs in the N-terminal region are the least conserved, while motifs in the central region are the most conserved (Fig. 3). Many of these motifs, such as the telomerase-specific “T” motif, are conserved only in telomerase RT, and thus serve as sequence signatures to distinguish telomerase RT from other RTs (Nakamura et al., 1997). The presence of these telomerase-specific signatures supports that the *Ciona* TERTs identified in this study are orthologs of vertebrate TERT. Furthermore, the phylogenetic relationship of the chordate TERT sequences reveals the evolutionary position of the identified *Ciona* TERT (Fig. 4), and is consistent with the phylogeny of eukaryotes based on sequences of other molecules, such as hemoglobin (Czelusniak et al., 1990).

### 3.3. Expression pattern of Ciona TERT mRNA

In humans, telomerase activity is usually regulated at the transcriptional level of the TERT gene. The expression of human TERT is restricted to those tissues of high turnover such as testis, intestine and bone marrow (Kolquist et al., 1998; Ulaner et al., 1998). To examine the pattern of TERT expression in *Ciona*, we examined the presence of the *Ciona* TERT transcript in a variety of tissues by RT-PCR. Two primer pairs were used to amplify the cDNA between the exons 5 and 15 or between the exons 15 and 17. PCR products of expected sizes were observed for both TERT and actin genes (Fig. 5). Alternative splicing products of TERT mRNA were not observed (data not shown). *Ciona* TERT mRNAs were detected in the intestine, stomach, ovary, muscle and testis (Fig. 5, lanes 3–7). The pattern is largely consistent with the known requirement of telomerase in tissues of high turnover, except the stomach and muscle (Fig. 5, lanes 4 and 6). In vertebrates, TERT is not expressed in stomach and muscle tissues (Yap et al., 2005). Since TERT is normally expressed in the tissues with significant regenerative capacity such as liver, one explanation would be that *Ciona* stomach and muscle tissue possess higher levels of regenerative capacity than that of human.

In normal human somatic cells, the TERT mRNA levels and telomerase activity are generally undetectable (Kim et al., 1994; Meyerson et al., 1997; Nakamura et al., 1997; Kolquist et al., 1996; Ulaner et al., 1998). In a teleost fish, *Fugu rubripes*, TERT is down-regulated in the brain and heart (Yap et al., 2005). Similarly, *Ciona* TERT mRNA also is not detectable in the brain and heart (Fig. 5, lanes 1 and 2). The specific down-regulation of the TERT gene in brain and heart tissue might make sea squirts an attractive model system for studying telomerase regulation during cell differentiation and organ development. However, in some organisms such as mouse, telomerase activity is not well regulated in adult tissues, and does not correlate with the expression of TERT mRNA (Martin-Rivera et al., 1998). To fully understand the regulation of telomerase in *Ciona*, both TERT protein level, telomerase RNA expression and telomerase activity will need to be measured.
from various tissues. This will then provide a foundation to a better understanding of the origin and evolution of telomerase regulation in vertebrates.

In conclusion, this work describes the identification and characterization of TERT protein from invertebrate chordates, C. intestinalis and C. savignyi. The Ciona telomerase provides an attractive system with an ideal evolutionary distance that is close enough to humans to identify commonality, but divergent enough to reveal conserved and critical features of the gene. To thoroughly study the biochemistry of Ciona telomerase, we would need to identify the telomerase RNA in Ciona. It would then allow us to reconstitute active Ciona telomerase enzyme for functional analysis. The cloning of Ciona TERT therefore offers a biochemical and structural basis for these future studies on Ciona telomerase.

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