Structure and function of echinoderm telomerase RNA

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ABSTRACT
Telomerase is a ribonucleoprotein (RNP) enzyme that requires an integral telomerase RNA (TR) subunit, in addition to the catalytic telomerase reverse transcriptase (TERT), for enzymatic function. The secondary structures of TRs from the three major groups of species, ciliates, fungi, and vertebrates, have been studied extensively and demonstrate dramatic diversity. Herein, we report the first comprehensive secondary structure of TR from echinoderms—marine invertebrates closely related to vertebrates—determined by phylogenetic comparative analysis of 16 TR sequences from three separate echinoderm classes. Similar to vertebrate TR, echinoderm TR contains the highly conserved template/pseudoknot and H/ACA domains. However, echinoderm TR lacks the ancestral CR4/5 structural domain found throughout vertebrate and fungal TRs. Instead, echinoderm TR contains a distinct simple helical region, termed eCR4/5, that is functionally equivalent to the CR4/5 domain. The urchin and brittle star eCR4/5 domains bind specifically to their respective TERT proteins and stimulate telomerase activity. Distinct from vertebrate telomerases, the echinoderm TR template/pseudoknot domain with the TERT protein is sufficient to reconstitute significant telomerase activity. This gain-of-function of the echinoderm template/pseudoknot domain for conferring telomerase activity presumably facilitated the rapid structural evolution of the eCR4/5 domain throughout the echinoderm lineage. Additionally, echinoderm TR utilizes the template-adjacent P1.1 helix as a physical template boundary element to prevent nontelomeric DNA synthesis, a mechanism used by ciliate and fungal TRs. Thus, the chimeric and eccentric structural features of echinoderm TR provide unparalleled insights into the rapid evolution of telomerase RNP structure and function.

Keywords: telomere; RNA structure; ribonucleoprotein; template boundary; evolution

INTRODUCTION
The telomerase ribonucleoprotein (RNP) enzyme synthesizes short DNA repeats onto linear eukaryotic chromosome termini to offset the progressive loss of telomeric DNA during replication, ensuring genome stability and cellular replicative capacity (Zakian 2009). The minimal telomerase enzyme necessary and sufficient for the reconstitution of telomerase activity is composed of the catalytic telomerase reverse transcriptase (TERT) and the telomerase RNA (TR) component that harbors a short template region for telomeric DNA synthesis (Podlevsky and Chen 2012). TRs from evolutionarily distinct groups of eukaryotic species are profoundly divergent, differing immensely in nucleotide sequence, length, and overall secondary structure (Chen and Greider 2004; Podlevsky et al. 2008).

Despite the massive difference in gross architecture, all known TRs contain two universal and indispensable structural domains: the template proximal pseudoknot and a stimulatory distal stem–loop moiety (Chen et al. 2002; Schmidt and Cech 2015). Within vertebrate TRs, this distal stem–loop moiety is termed CR4/5 (conserved regions 4 and 5) and composed of a three-way junction of the P5, P6, and P6.1 stem–loops. A remarkably similar structure with an identical function has been recently identified and found to be absolutely conserved across the evolutionary distant filamentous fungal and fission yeast TRs (Qi et al. 2013). In addition to the template/pseudoknot and CR4/5 domains, there are immense species-specific structural differences within TR that appear in concert with the plethora of species-specific TR binding proteins. This includes RNA helicase RHAU, TCAB1, and box H/ACA dyskerin protein complex in vertebrates; Est1, Est3, Ku70/80 heterodimer, and Sm ring complex in fungi; p50 and p65 in ciliates; and box C/D Nop58 and MTAP (TCAB1 homolog) in flagellates (Mitchell et al. 1999; Seto et al. 1999; Hughes et al. 2000; Ventecher et al. 2009; Lattmann et al. 2011; Sexton and Collins 2011; Egan and Collins 2012; Singh et al. 2012; Tang et al. 2012; Gupta et al. 2013). These unique sets of TR accessory

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proteins are essential for TR localization, maturation, and RNP assembly (Schmidt and Cech 2015).

Due to the close proximity of echinoderms to vertebrates, the echinoderm TR identified from the purple sea urchin (Strongylocentrotus purpuratus) has the expected conservation of the template proximal pseudoknot and the 3′ H/ACA domains, homologous to those in vertebrate TR (Li et al. 2013). However, it was surprising that the central domain of purple sea urchin TR would lack the ancestral vertebrate CR4/5 structural domain. Thus, additional sequences and biochemical analysis of echinoderm TR are necessary to determine the extent of conservation for these apparent echinoderm-specific structural elements as well as their functional relevance for telomerase RNP assembly and enzymatic activity.

To determine echinoderm TR secondary structure by phylogenetic sequence analysis, it is necessary to obtain TR sequences from each major clade of echinoderm species. However, this task is complicated by the high variation of TR sequences from even closely related groups of species, which prevents degenerate PCR-based strategies (Chen et al. 2000; Podlevsky and Chen 2012). A variety of biochemical enrichment protocols and computational screening methodologies have been developed to improve the rate of TR identification from previously unexplored groups of species (Greider and Blackburn 1989; Leonardi et al. 2008; Webb and Zakian 2008; Xie et al. 2008; Gunisova et al. 2009; Cifuentes-Rojas et al. 2011; Li et al. 2013; Qi et al. 2013). Unfortunately, the vast majority of these methodologies are incompatible with TR identification from echinoderm species; as they typically rely on either established genetic tools, small genome size, abundant telomerase enzyme, or a longer template length. Furthermore, with the exception of the purple sea urchin, no echinoderm species has a complete genome sequenced, assembled, annotated and publicly available (Sodergren et al. 2006; Cameron et al. 2015). The scarcity of completed and annotated echinoderm genomes precludes the approach of targeting syntenic protein genes flanking the TR gene, which has been successfully used for several sensu stricto Saccharomyces species (Dandjinou et al. 2004) and plants (Beilstein et al. 2012). Thus, ancillary avenues for the identification of additional echinoderm TRs are necessary to elucidate the exceedingly divergent TR structures from this group of species.

Herein we report the secondary structure of echinoderm TR determined by phylogenetic comparative and biochemical analyses. Similar to the purple sea urchin TR, additional echinoderm TRs identified in this study contain a conserved template/pseudoknot domain with a template-adjacent helix for template boundary definition and an H/ACA domain for biogenesis. The structure of the purple sea urchin TR central domain is moderately conserved in other sea urchin and sand dollar species, while brittle and feather star TRs have a distinct structure in their central domain. Functional analysis revealed that the echinoderm template/pseudoknot domain alone is sufficient to reconstitute significant telomerase activity with the TERT protein, while a short helical region from the central domain moderately stimulates activity. The chimeric structural features of echinoderm TR demonstrate the innate molecular flexibility of the telomerase RNP for maintaining sufficient function yet permitting rapid divergence in sequence and structure of integral TR domains.

RESULTS

The identification of the first invertebrate TR from the purple sea urchin provided an initial glimpse of conservation and the peculiar loss of critical vertebrate TR structural features within echinoderms (Li et al. 2013). Despite the close evolutionary relationship between vertebrates and echinoderms, the essential CR4/5 structural domain of vertebrate TR is seemingly absent from purple sea urchin TR. This was unexpected since the CR4/5 domain is absolutely conserved in the evolutionarily distant vertebrate, filamentous fungal and fission yeast TRs (Qi et al. 2013). To elucidate the extent to which ancestral structural features of TR have diversified and become unrecognizable within echinoderms, while retaining telomerase function, we identified 14 new invertebrate TRs from a broad range of echinoderm species using three distinct strategies and performed phylogenetic comparative analysis to infer the conserved secondary structure of echinoderm TR.

The strategies used in this study for TR identification are different from our previous approach for identification of purple sea urchin TR (Li et al. 2013). In our previous study, the biochemical enrichment of purple sea urchin TR relied on immunoprecipitation using antibodies targeting the 5′-TMG cap. While a 5′-TMG cap has been experimentally confirmed for vertebrate, yeast, and the purple sea urchin TRs, very little is known for other echinoderm TRs (Seto et al. 1999; Jády et al. 2004; Li et al. 2013). The additional TR biochemical enrichment step depended on immunoprecipitation with the corresponding recombinant-tagged TERT protein. This step is challenging due to the lack of additional echinoderm TERT proteins identified (Podlevsky et al. 2008). Moreover, the lack of completed echinoderm genomes impedes echinoderm TERT protein identification and precludes syntenic gene searches (Sodergren et al. 2006; Cameron et al. 2015). To overcome these challenges, we therefore developed a separate TR biochemical enrichment strategy for identifying TR from additional echinoderm species, avoiding the anti-TMG and TERT-binding immunoprecipitation steps.

Our initial strategy for echinoderm TR identification from species closely related to purple sea urchin was degenerate PCR with primers targeting sequences within the template/pseudoknot and H/ACA domains (Chen et al. 2000; Li et al. 2013). The targeted sequences are well conserved between purple sea urchin and vertebrate TRs. With this approach, we successfully identified three new echinoderm
Identification of echinoderm TRs. Eight sea urchins, a sand dollar (Dendraster excentricus), three brittle star, and a feather star (Anniesia japonica) TR species were identified in this study (black text) with two sea urchin TRs (violet text) previously identified (Li et al. 2013; Gillard et al. 2014). The TR length was determined by 5′- and 3′-RACE, estimated from the position of the box ACA in the sequencing data (*), or not determined (n/d). Echinoderms comprise the sister classes Echinoidea (sea urchins and sand dollars), Ophiuroidea (brittle stars), and the ancestral Crinoidea class (sea lilies and feather stars). The vertebrate lineage (phylum Chordata, blue) is closely related to the echinoderm lineage (light violet), while fungi (phylum Ascomycota, green) are the out-group.

TR sequences from two distinct orders of sea urchins: the purple-spined sea urchin (Arbacia punctulata) from the order Arbacoidea as well as the green variegated (Lytechinus variegatus), and blue tuxedo (Mespilia globulus) sea urchins from the order Temnopleuroidea (Fig. 1). Sequence alignment of the identified sea urchin TRs revealed a highly conserved sequence in the template/pseudoknot domain, similar to that of vertebrate TRs (Supplemental Fig. S1). However, further attempts to identify TRs from the more distantly related echinoderm orders using a degenerate PCR strategy were unsuccessful and necessitated an alternative approach.

To identify TRs from more distantly related echinoderm orders, we used a next-generation sequencing-based approach adapted from our previously reported strategy (Li et al. 2013). We chose two target species outside of the sea urchin family: the eccentric sand dollar (Dendraster excentricus) and Panamanian serpent brittle star (Ophioderma panamense) (Fig. 1). Sand dollars are closely related to sea urchins, belonging to the same echinoderm class Echinoidea, yet from a separate order Clypeasteroida (Pereseke et al. 2010). Brittle stars are more distantly related, belonging to the separate class Ophiuroidea. Total RNA was isolated from gonadal tissues and enriched for TR by a size-selection step to deplete ribosomal RNAs, mRNAs and small RNAs prior to RNA-seq analysis. RNA species ranging in size from 300 to 700 nt were purified by polyacrylamide gel electrophoresis (PAGE). This size range was chosen based on the length of the four sea urchin TRs (530–563 nt) and all known vertebrate TRs (312–559 nt) (Podlevsky et al. 2008). A caveat of this purification step was that the TR of interest must have a length within this range. The size-selected RNA was then used for cDNA library construction (see Materials and Methods). Next-generation sequencing of the two cDNA libraries generated over 75 million and 77 million 50-bp sequencing reads for sand dollar and brittle star, respectively. The RNA-seq data from these two species were independently de novo assembled into contigs using the Trinity assembly program (Haas et al. 2013). The assembled contigs were searched for putative TR sequences using the Fragrep2 program (Mosig et al. 2006) and position-specific weight matrices (PWM) derived from the sequence alignment of the four sea urchin TRs (Supplemental Fig. S1B). The Fragrep2 search produced a single hit from each of the sand dollar and brittle star data sets. Mapping the respective TR 5′- and 3′-ends by RACE revealed that the eccentric sand dollar and brittle star TRs are 558 and 427 nt in length, respectively, within the range of our PAGE size selection (Fig. 1). Each TR demonstrated sequence conservation in the template/pseudoknot and H/ACA domains, however, the brittle star TR central domain could not be aligned with urchin TR sequences (Supplemental Figs. S2, S3).

For our last echinoderm TR search strategy, we then performed computational analysis of publicly available transcriptome data. The echinoderm transcriptome data obtained from the National Center for Biotechnology Information Sequencing Read Archive (http://www.ncbi.nlm.nih.gov/sra) were de novo assembled into contigs and searched with the Fragrep2 program using a new PWM pattern improved by including the sand dollar and brittle star TR sequences. Through this computational approach, we identified nine TR sequences from three separate echinoderm classes: five sea urchins from Echinoidea, two brittle stars from Ophiuroidea, and a feather star from the basal class Crinoidea (Fig. 1).

**Echinoderm TR utilizes a distinct mechanism for template boundary definition**

The secondary structures of echinoderm TRs from purple sea urchin, blunt spined brittle star, and feather star—each representing a distinct echinoderm class—were discerned by phylogenetic comparative analysis (Fig. 1). Phylogenetic comparative analysis identifies nucleotides that vary in concert among species while maintaining Watson–Crick base-pairing, inferring helical regions in the secondary structure (Chen et al. 2000). Due to the low level of sequence conservation between echinoderm classes, TR primary sequences from each class of species were aligned separately (Supplemental Figs. S2–S4). The secondary structures for the three echinoderm TRs were constructed based on nucleotide covariations from these multiple sequence alignments.
Despite the limited number of universally conserved residues, the template/pseudoknot domain of echinoderm and vertebrate TRs share homologous architectures (Fig. 2A–D). Interestingly, the echinoderm TR template/pseudoknot domain harbors a short template-adjacent helix, termed P1.1, downstream from the pseudoknot core enclosing P1 stem. While a template-adjacent stem–loop is commonly present in fungal and specific ciliate TRs, it is absent in all known vertebrate

FIGURE 2. Echinoderm TR template/pseudoknot domain contains a template-adjacent helix for template boundary definition. Comparison of representative TR template/pseudoknot domains from vertebrate, human (A); sea urchin, purple sea urchin (B); brittle star, blunt spined brittle star (C); and feather star, Japanese feather star (D). The hallmark triple helix within the TR pseudoknot (green) is denoted within each structure. Covariation (black bar), universal-invariant between vertebrates and echinoderm (red), and group-invariant (orange) nucleotides for vertebrate, sea urchin, and brittle star TR template/pseudoknot domains are based on multiple sequence alignment of 42 vertebrate (Chen et al. 2000; Podlevsky et al. 2008; Xie et al. 2008), nine sea urchin/sand dollar, three brittle star, and a feather star species (see Supplemental Fig. S2). (E–G, top) Schematic of the purple sea urchin TR template/pseudoknot domain denoting nucleotide mutations (violet), insertions (green), and truncations to determine P1.1 functionality for template definition. (E–G, bottom) Functional analysis of purple sea urchin TR mutations and truncation variants by the direct primer-extension assay. Sea urchin TR template/pseudoknot variants (fragment 11–163 nt) and the central domain fragment (186–456 nt) were assembled in vitro with the sea urchin TERT protein. Sea urchin telomerase variants were assayed with the 18-mer DNA primer (TTAGGG), and this primer 32P end-labeled was added as a loading control (l.c.). The incorporation of nontelomeric nucleotides from the downstream-flanking region of the template, read-through, is denoted on the gel (red triangles).
TRs (Chen et al. 2000; Lai et al. 2002; Dandjinou et al. 2004). Nonetheless, one echinoderm TR from the spiny brittle star (*Ophiothrix spiculata*) appears to lack a P1.1 stem–loop and structurally better resembles the vertebrate template/pseudoknot domain (Supplemental Fig. S2A).

In vertebrate TR, the template boundary is defined by the core-enclosing P1b helix that restricts the availability of residues to be used as templates (Chen and Greider 2003). In contrast, the template boundary in fungal TR is defined by the template-adjacent helix (Tzfati et al. 2000; Dandjinou et al. 2004; Leonardi et al. 2008; Webb and Zakian 2008). To discern whether the echinoderm template-adjacent P1.1 helix functions as a template boundary element similar to fungal TR, we reconstituted purple sea urchin telomerase in vitro with TR mutants bearing insertions and/or deletions intended to alter the structure or shift the position of the P1.1 helix (Fig. 2E, top). To monitor nucleotide incorporation bypassing the template boundary, we generated the sea urchin TR mutant C22U/U41G in which the mutated 41G residue can be used as the template in the presence of dCTP when nucleotide addition bypasses the template boundary. The mutated residue 22U was introduced to prevent unintended base-pairing between the native C22 residue and 41G, which could potentially extend the P1.1 helix and limit the use of 41G as template. Using the C22U/U41G mutant as the parental construct, we generated three mutants, linker-1, -2, and -3, with insertions or deletions in the regions flanking the P1.1 helix (Fig. 2E, top). These telomerase mutants were reconstituted in vitro and analyzed by the direct primer-extension assay to assess enzymatic activity and template boundary definition. Our results showed that the linker-1 mutant with the insertion of two uridine residues between the template and the P1.1 stem–loop used 41G as template for nucleotide incorporation in the presence of dCTP, bypassing the normal template boundary (Fig. 2E, bottom, lanes 5,6). In vertebrate TR, the length of the linker between the template and stem P1b is critical for template boundary definition (Chen and Greider 2003). We thus tested the importance of the distance between the template and P1b in purple sea urchin TR for maintaining template boundary. We generated mutant linker-2 that had the same two-residue insertion as the linker-1 mutant and a two-residue deletion in the linker between the P1b and P1.1 helices (Fig. 2E, top). The combination of the insertion and deletion maintained the overall distance between the template and P1b (Fig. 2E, top). Interestingly, the linker-2 mutation failed to prevent nucleotide addition bypassing the template boundary (Fig. 2E, bottom, lanes 7,8). This suggests that the linker length between P1b and P1.1, critical for vertebrate TR template boundary definition, is not important for template boundary definition in purple sea urchin. Furthermore, the linker-3 mutant, with two residues inserted in the linker between P1b and P1.1, had no significant effect on template boundary (Fig. 2E, lanes 9,10). Therefore, the P1.1 stem–loop is crucial for defining the echinoderm TR template boundary.

We then examined whether the echinoderm TR P1b can function as the template boundary element when the P1.1 helix has been removed. To test this, we first confirmed that the sea urchin P1b helix was dispensable in the presence of the P1.1 helix by removing the entire P1b helix with mutant ΔP1b (Fig. 2F, top). The loss of the P1b helix did not disrupt the template boundary; the addition of dCTP did not alter the banding pattern of telomerase products (Fig. 2F, bottom, lanes 5,6). To explore the possibility of converting the sea urchin TR to use a vertebrate template boundary element, we generated mutants ΔP1.1-1, -2, -3, and -4 by removing the P1.1 helix and introducing serial single nucleotide deletions in the new linker between the template and P1b (Fig. 2F, top). Our results demonstrated that mutants ΔP1.1-1 and -2 with longer linker lengths between the template and P1b permitted visible nucleotide addition bypassing the end of the template in the presence of dCTP (Fig. 2F, bottom, lanes 7–10). Interestingly, mutants ΔP1.1-3 and -4, with shorter linker lengths between the template and P1b, showed little to no nucleotide addition bypassing the template boundary in the presence of dCTP (Fig. 2F, bottom, lanes 11–14). Thus, in the absence of P1.1, the length of the linker between the template and P1b appeared critical for template boundary definition, similar to the vertebrate mechanism.

We further investigated whether the helical structure of P1.1 is important for template boundary definition. Mutants m1 and m2 harbored mutations in one of the two strands to disrupt base-pairings, while mutant m3 had compensatory mutations to restore base-pairings in the P1.1 helix (Fig. 2G, top). The m1 and m2 mutant showed impaired template boundary definition, allowing incorporation of dCTP using 41G as the template (Fig. 2G, bottom, lanes 6–8). The m3 mutant prevented nucleotide addition bypassing the end of the template in the absence or presence of dCTP, restoring the template boundary (Fig. 2G, bottom, lanes 9,10). Together, these data indicated that the P1.1 stem–loop, and not the P1b helix, functions as the template boundary element in echinoderm TR. However, the loss of the P1.1 helix can be compensated for by the P1b helix with a proper linker length, functioning as an ancillary template boundary.

**The echinoderm CR4/5 structural element**

Sea urchin, brittle star, and feather star TRs share no apparent sequence or structural homology in the central region, which lies between the template/pseudoknot and H/ACA domains. Moreover, the primary sequences of the echinodermal central region could not be aligned with the conserved CR4/5 sequences of vertebrate TR. We therefore discerned the secondary structures of the echinoderm central domain individually for each of the three echinoderm classes (Supplemental Fig. S3A,C). For sea urchin TRs, the alignment of the central domain sequences revealed noticeable sequence conservation within the latter half of this region (Supplemental Fig. S3A). To construct a secondary structure model of the sea urchin
TR central domain, the aligned sequences were analyzed for nucleotide covariation. Secondary structure prediction was assisted by the mFold program (Zuker 2003). A preliminary model predicted two separate long stem–loop branches connected at their bases by the conserved P4 helix (Fig. 3B). The 3′-proximal helix, termed P4.2, showed a significantly higher sequence conservation than the 5′-proximal P4.1 helix (Supplemental Fig. S3A). For the brittle star TR, the central domain was predicted to fold into a single helix (Fig. 3C; Supplemental Fig. S3C). The small number of brittle star TRs did not provide appreciable nucleotide covariation support. The feather star TR central domain was also predicted to form a single, while much shorter, helix that is similar to the brittle star central domain (Fig. 3D). The identification of additional TRs from brittle and feather star species would be necessary to lend further nucleotide covariation support to these predicted single helical structures.

To refine the secondary structure of echinoderm TR central domain, we performed selective 2′-hydroxyl acylation analyzed by primer extension (SHAPE) to probe for unpaired nucleotides in the RNA structure. Overall, there was considerable agreement between the SHAPE data and the secondary structure derived from the sequence alignment of 10 sea urchin TR sequences (Fig. 3B; Supplemental Fig. S3A,B). For the brittle star TR, the SHAPE analysis supported the central domain as a single helical structure (Fig. 3C; Supplemental Fig. S3C,D). Therefore, the secondary structure of the echinoderm central domain is unique among TRs, lacking the three-way junction configuration of the essential and ancestral CR4/5 structure found in vertebrate and fungal TRs (Chen et al. 2002; Qi et al. 2013).

Functional analysis of the echinoderm TR central domain was performed to determine the minimal region necessary and sufficient for reconstituting telomerase enzymatic activity (Fig. 4; Supplemental Fig. S4). We previously found that the purple sea urchin template/pseudoknot TR fragment and TERT protein are sufficient to reconstitute a basal level of telomerase activity, ∼40% of that generated with the inclusion of the P4.2 helix (Li et al. 2013). To further identify the regions in this P4.2 helical fragment dispensable for telomerase activity stimulation, we generated purple sea urchin TR P4.2 helix fragments, P4.2-Δ1, -Δ2, and -Δ3, by serially truncating from the apical loop and capping with the tetra-loop GAAA (Fig. 4A). Purple sea urchin telomerase was reconstituted with the purple sea urchin TERT, the template/pseudoknot domain, and these various P4.2 truncation fragments (Fig. 4B). Truncation mutation P4.2-Δ1, which removed nucleotides 342–388 from the helix, significantly reduced telomerase activity to 49% of the activity reconstituted with the entire P4.2 helix (Fig. 4B, lanes 1,4). The reduced activity of mutant P4.2-Δ1 was similar to the basal activity of the telomerase lacking the P4.2 helix (Fig. 4B, lanes 2,3). In contrast, truncation mutations P4.2-Δ2 and -Δ3 that removed the apical loop and helical regions from nucleotides 350–379 and 358–373, respectively, did not reduce telomerase activity (Fig. 4B, lanes 5,6). This suggests that the region from nucleotides 350–379 nt is not required for the function of the P4.2 helix.

We previously mapped the region of the P4.2 helix essential for activity stimulation to between nucleotides 332 and 400 (Li et al. 2013). By combining the previous minimal fragment with the P4.2-Δ2 truncation, we generated a 43-nt P4.2 fragment covering nucleotides 332–349 and nucleotides 380–400, termed P4.2-Δ2-mini (Fig. 4C). As expected, this small RNA fragment was sufficient to stimulate telomerase activity to a similar level as the entire central P4.2 fragment (Fig. 4D, lane 4). Therefore, despite the considerable differences in the primary sequences and secondary structures, we concluded that the P4.2-Δ2-mini helix contains the structural element functionally equivalent to the vertebrate CR4/5 domain necessary for stimulating telomerase activity. In recognition of this functional homology, we termed the helical structure of P4.2-Δ2-mini the eCR4/5 domain for functionally equivalent to the vertebrate CR4/5 domain.

The eCR4/5 domain of purple sea urchin TR contains two bulged residues, C345 and U392, that are conserved across
sea urchin and sand dollar species (Fig. 3B; Supplemental Fig. S3A). Within the vertebrate CR4/5 domain, the P6 stem–loop contains a conserved single-nucleotide uridine bulge (Chen et al. 2000; Xie et al. 2008). The deletion of this uridine bulge drastically reduces binding of the medaka fish CR4/5 domain to its TERT protein (Bley et al. 2011). To investigate whether either single-nucleotide bulge in purple sea urchin eCR4/5 has a similar vital function for RNA–protein interaction, we generated single-residue deletion mutants of P4.2-Δ2-mini: ΔC345 and ΔU392 (Fig. 4C). The individual deletion of C345 or U392 abolished or severely reduced the stimulatory function of eCR4/5 (Fig. 4E, lanes 2–4). Additionally, the telomerase stimulatory effect of echinoderm eCR4/5 is species-specific, as the vertebrate CR4/5 domain fragment from medaka fish TR failed to enhance activity of purple sea urchin telomerase (Fig. 4E, lane 5).

We further expanded our functional analysis of the echinoderm central domain to include the blunt spined brittle star (Supplemental Fig. S4). We generated various central domain fragments, CDΔ1, -Δ2, -Δ3, -Δ4, and -Δ5, of the blunt spined brittle star TR by serially truncating from the base of the helix retained similar telomerase activity as the entire central domain, demonstrating that the regions removed are dispensable (Supplemental Fig. S4B, lanes 1–5). In contrast, truncation mutation CDΔ5 reduced activity to 67%, similar to the basal level of activity generated with the template/pseudoknot domain alone, indicating the region removed from the CD-Δ4 fragment is crucial for activity (Supplemental Fig. S4B, lanes 6,7). We then generated additional truncation mutants, ALΔ1, -Δ2, -Δ3, -Δ4, and -Δ5, by serially truncating the brittle star TR central domain from the apical loop and capping the stem with the tetra-loop GAAA (Supplemental Fig. S4C). The activity assay of mutants ALΔ1 showed that the apical loop of brittle star TR central domain is dispensable for the stimulation of telomerase activity (Supplemental Fig. S4D, lanes 1,2). However, additional truncations from the apical loop with mutants ALΔ2, -Δ3, -Δ4, and -Δ5 eliminated the stimulatory effect of the central domain fragment (Supplemental Fig. S4D, lanes 3–7). Thus, the brittle star eCR4/5 domain, like that of sea urchin, is an internalized helical region as opposed to the three-way-junction of helices found in the vertebrate CR4/5 domain. Interestingly, the brittle star reconstituted a basal level of telomerase activity at ∼70% of the full activity, while the sea urchin template/pseudoknot-TERT RNP generated only 40% of the full activity. Together, these results suggest that the sea urchin and brittle star eCR4/5 are functionally equivalent to the vertebrate
CR4/5 despite the lack of sequence or structural conservation (Fig. 3).

**Echinoderm H/ACA domain**

The 3’ region of echinoderm TR harbors an H/ACA domain with the highly conserved sequence motifs and a secondary structure homologous to that of vertebrate TR (Fig. 5). Similar to the vertebrate H/ACA domain, the echinoderm H/ACA domain comprises the two universally conserved box H and ACA motifs separated by a stem–loop with two paired regions, P7 and P8. RACE analysis for select echinoderm species determined that the 3’-end of echinoderm TR lies three nucleotides downstream from the box ACA motif, identical to the vertebrate TR 3’-end (Chen et al. 2000; Xie et al. 2008). In sea urchin and brittle star TRs, the apical loop capping the P8 helix contains a conserved CAB box motif that is critical for vertebrate TR accumulation and Cajal body localization (Fig. 5B,C; Supplemental Fig. S5). However, a CAB box could not be found in the P8 loop of feather star TR (Fig. 5D). This apparent loss of the CAB box within feather star in echinoderms is not overly surprising as teleost fish TR also lacks a CAB box (Xie et al. 2008). The conservation of the H/ACA domain in echinoderm TRs suggested that the H/ACA small nucleolar/small Cajal body (sno-/sca-) RNA biogenesis pathway and 3’-end processing mechanism are similarly used by both vertebrate and echinoderm TRs.

**DISCUSSION**

The initial identification of purple sea urchin TR revealed structural domains homologous to the template/pseudoknot and H/ACA domains from vertebrate TRs (Li et al. 2013). Surprisingly, purple sea urchin TR seemingly lacks the essential ancestral CR4/5 structural domain that is evolutionarily conserved and functionally essential for vertebrate, filamentous fungal, and fission yeast telomerase activity (Qi et al. 2013). Phylogenetic comparative analysis of 14 echinoderm TRs, identified in this study, indicates that the CR4/5 three-way junction of helices has been functionally replaced with a simple internal helical region (Fig. 4). Interestingly, the overall architecture of echinoderm TRs appears chimeric, with elements common to vertebrate or fungal TRs, as well as echinoderm-specific structural features (Fig. 6).

The strategies that we used in this study to identify TR sequences from the echinoderm species rely on powerful RNA-seq analysis of size-selected RNA samples and de novo transcriptome assembly of the sequencing reads. In addition, our simplified bioinformatics screening step using Fregrep2 program generated PWM facilitated identification of more divergent TR sequences than our previously used TR-PK-finder algorithm (Supplemental Fig. S1B). This spartan approach was successful for echinoderm TR identification and can be readily applied toward the identification of TRs from other unexplored groups of species that lack sequenced genomes and biochemically characterized telomerase.

Our functional characterization of the purple sea urchin TR template/pseudoknot domain revealed that the echinoderm TR template boundary is defined by the template-adjacent P1.1 stem–loop (Fig. 2). The P1.1 stem–loop resembles the template-adjacent helix in fungal TRs and Helix II in ciliate TRs (Podlevsky and Chen 2012). Fungal and ciliate template-adjacent helices function similarly by physically restricting the residues in the template-flanking region from being used as the template (Tzafiti et al. 2000; Jansson et al. 2015). Instead of using a template-adjacent helix, vertebrate TR template boundary is defined by the more distant P1b helix and a single-stranded linker with a specific length to the template (Chen and Greider 2003). Due to its evolutionary proximity to vertebrates, echinoderm TR was expected to have a template boundary element reminiscent of the vertebrate mechanism (Fig. 1). It is rather surprising that echinoderm TR utilizes a template-adjacent helix to define its template boundary, a mechanism that is highly similar to fungal and ciliate.
TRs (Fig. 2). However, it remains unclear whether the echinoderm P1.1 stem-loop functions as a TERT protein binding site, similar to ciliate TR helix II, whereby TERT protein binding impedes the use of nucleotides as the template from outside the template boundary element (Autexier and Greider 1995; Lai et al. 2002; Jansson et al. 2015). Moreover, as demonstrated in this study, the echinoderm P1.1 helix can be entirely removed and replaced with a specific linker length to the P1b helix which functions as the new template boundary element, using a mechanism identical to that of vertebrate TRs (Fig. 2F). Interestingly, the echinoderm P1.1 helix appears to have been lost within spiny brittle star TR, yet present within other brittle star TRs (Supplemental Fig. S2A). However, we suspect that the loss of P1.1 in spiny brittle star TR was an isolated event during evolution as TRs from other brittle stars and the more basal echinoderm species, feather star, preserve the template-adjacent P1.1 helix (Fig. 2).

The conserved three-way junction structure of the CR4/5 domain has been replaced with a simple helical structural eCR4/5 domain in echinoderm TR (Fig. 3). The echinoderm eCR4/5 domain is functionally equivalent to the vertebrate and fungal CR4/5 domain, which is to stimulate telomerase activity (Fig. 4; Supplemental Fig. S4). The transition from the CR4/5 to the eCR4/5 domain along the echinoderm lineage was likely prompted by the gain-of-function of the echinoderm template/pseudoknot domain. Significant telomerase activity can be reconstituted from the echinoderm TERT and TR template/pseudoknot domain alone (Fig. 4B; Supplemental Fig. S4), a sharp contrast to vertebrate and filamentous fungal template/pseudoknot-TERT RNP complexes that rely absolutely upon CR4/5 binding for detectable telomerase activity (Chen et al. 2002; Qi et al. 2013). Due to the ancestry of the CR4/5 structural domain, it would appear that this critical and evolutionarily conserved structural element has diversified within the echinoderm phylum of species. The apparent loss of functional dependency on the CR4/5 is not exclusive to echinoderm TR. The budding yeast TR, while having preserved the three-way junction of helices, lacks the highly conserved P6.1 stem-loop (Brown et al. 2007). Furthermore, a miniature version of the budding yeast TR that contains the template/pseudoknot domain, and lacks the three-way junction, is sufficient to confer a detectable telomerase activity and maintain telomere length (Zappulla et al. 2005). This suggests that the loss of function of the CR4/5 domain has been compensated by the gain of function of the yeast template/pseudoknot domain.

Throughout the echinoderm lineage, TR contains a 3′ H/ACA domain that has a highly conserved secondary structure and motifs homologous to the vertebrate H/ACA domain (Fig. 5). This strongly supports a similar TR sno-scaRNA biogenesis pathway and 3′-end processing mechanism between vertebrate and echinoderm TRs (Jády et al. 2004; Egan and Collins 2010). Additionally, the presence of a CAB box in the apical loop of the P8 helix in the echinoderm H/ACA domain indicates binding by a homolog of the vertebrate TCAB1 protein for Cajal body localization (Ventechier et al. 2009). Unexpectedly, the feather star H/ACA domain lacks a CAB box in the P8 apical loop, which is similar to teleost fish TR as a vertebrate outlier (Xie et al. 2008). The absence of an apparent CAB box would indicate either cryptic binding by a TCAB1 homolog, an alternative means of Cajal body localization, or an alternative biogenesis pathway.

The structural divergence of the eCR4/5 domain within various echinoderm TRs and the gain-of-function of the template/pseudoknot domain demonstrate functional coevolution of these two separate TR structural domains. Echinoderm template/pseudoknot-TERT RNP confers significant telomerase activity, reducing telomerase dependence on the ancestral CR4/5 domain. This would have permitted the rapid evolution and divergence in RNA structure by either gradual accumulation of mutations or a more drastic event that devastated and dismantled CR4/5 physically and functionally. Echinoderm TR has a template-adjacent P1.1 stem-loop that is reminiscent of, and functionally homologous
to, fungal and ciliate telomerases. Thus echinoderm TR possesses chimeric structural features—an amalgam of vertebrate and fungal functional elements, which expands our understanding of telomerase divergence and of RNA–protein coevolution.

MATERIALS AND METHODS

Isolation of total RNA

Gonadal tissue was dissected from live D. excentricus, O. panamense, and O. echinata (Marine Research and Educational Product and Live Aquaria). Total RNA was isolated from gonads using TRI-Reagent (Molecular Research Center, Inc.) following the manufacturer’s instructions appended with an acid phenol extraction step prior to chloroform extraction and ethanol precipitation. RNA quality was determined by electrophoresis on a 1% agarose/formaldehyde denaturing gel.

RNA preparation and RNA-seq

Ten micrograms of total RNA for D. excentricus and O. panamense was resolved by electrophoresis on a 4% polyacrylamide/8 M urea denaturing gel. The gel section, containing RNA species from 300–750 nt in length, was excised and RNA was eluted from the gel slice followed by ethanol precipitation. Size-selected RNA was used for cDNA library construction with the ScriptSeq v2 RNA-Seq Library Preparation Kit (Epicentre) following the manufacturer’s instructions. The cDNA libraries were constructed with ScriptSeq Index PCR Primers (Epicentre) and indexed cDNA libraries were pooled for a single multiplexed single-end 50-bp sequencing run on a HiSeq 2000 (Illumina).

Degenerate PCR

Gonadal tissue was dissected from live A. punctulata, L. variegatus, and M. globulus (Live Aquaria). Partial TR genes were amplified from genomic DNA extracted from gonadal tissues by PCR using degenerate DNA primers (Supplemental Table 1). A 12.5 µL PCR reaction was performed in 1× Ex Taq buffer (Takara) with 1 µM forward primer, 1 µM reverse primer, 100 µM of each dNTP, 100 ng of genomic DNA, and 0.3 U ExTaq Hot Start DNA polymerase (Clontech). Forward and reverse primers were designed to target conserved sequences in the pseudoknot and H/ACA domains, respectively. The full-length genomic sequence of each TR gene was determined by ligation-mediated PCR using the Universal Genome Walker Kit (Clontech) followed by sequencing as described previously (Chen et al. 2000).

Sequence alignment analysis

The multiple sequence alignment of vertebrate and echinoderm TRs was performed initially using the program BioEdit and the ClustalW algorithm. The alignments were further refined manually using the highly conserved regions and known motifs as anchor points. Closely related species were first aligned and the alignment expanded to include sequences from more divergent species.

Bioinformatics analysis of next-generation sequencing data

The next-generation sequencing data obtained from the pooled cDNA library samples were de-multiplexed and de novo assembled using the Trinity assembly program (Haas et al. 2013) with default parameters on the National Science Foundation Data Intensive Academic Grid shared computational cloud. The assembled transcripts were searched in the sense-strand direction using the Fragrep2 program (Mosig et al. 2006) with a PWM sequence pattern generated from the multiple sequence alignment of six vertebrate and four sea urchin TRs (Supplemental Fig. S1). This PWM pattern was relaxed for greater range in the spans between conserved sequences, permitting a more degenerate search. Transcriptome and genome data from NCBI SRA (http://www.ncbi.nlm.nih.gov/sra) were de novo assembled using the Trinity assembly program and searched with the Fragrep2 program.

Determination of RNA 5′- and 3′-ends

The 5′- and 3′-ends of D. excentricus, O. panamense, and O. echinata TRs were determined by Rapid Amplification of cDNA Ends (RACE) using the FirstChoice RLM-RACE kit (Ambion). For all other echinoderm TRs identified in this study, the 3′-end was predicted as 3 nt downstream from the box ACA motif as previously described (Chen et al. 2000).

SHAPE analysis

The central domain of O. echinata and S. purpuratus TR were PCR amplified with primers appended with SHAPE-specific adapter sequences, as previously described (Wilkinson et al. 2006). These PCR products were used as templates for in vitro transcription. The adapter appended RNA was gel purified and ethanol precipitated. Two picomoles of purified RNA fragments was denatured in 0.5× TE buffer (pH 8.0) at 95°C for 2 min and immediately placed on ice for 2 min. The RNA was supplemented with a final concentration 1× RNA folding mix (100 mM HEPES at pH 8.0, 6 mM MgCl₂, and 100 mM NaCl), incubated at 30°C for 20 min, and divided into half. The RNA was treated with either 6.5 mM N-methylisatoic anhydride (NMIA) (Sigma) in anhydrous DMSO or DMSO alone, incubated at 30°C for 1 h and 25 min (five NMIA hydrolysis half-lives), and ethanol precipitated. One picomole of 32P end-labeled primer was added to the RNA samples in 0.5× TE buffer (pH 8.0), incubated at 65°C for 5 min and 35°C for 5 min, and then placed on ice. The mixture was supplemented with 1× SuperScript III First-strand buffer (50 mM Tris-HCl at pH 8.3, 75 mM KCl, and 3 mM MgCl₂), 5 mM DTT, 0.5 mM each dNTP, and 0.75 mM ddGTP or ddATP for the untreated RNA, followed by incubation at 52°C for 1 min, addition of 100 U SuperScript III RT (Life Technologies), and incubation at 52°C for 10 min. Alkaline hydrolysis was performed with a final concentration of 200 mM sodium hydroxide and incubation at 95°C for 5 min and stopped by the addition of Acid Stop solution (77 mM unbuffered Tris-HCl, 32% formamide and 8 mM EDTA) incubated at 95°C for 5 min. The DNA products were resolved on a 6.8% polyacrylamide/8 M urea denaturing gel. The gel was dried, exposed to a phosphorstorage screen, and imaged on a phosphorimager FX-Pro (Bio-Rad). NMIA reactivity was normalized by subtracting
the lowest intensity value from all positions, followed by subtracting the intensity values of the DMSO control from the corresponding NMIA reaction.

Brittle star TERT gene synthesis

The brittle star (O. echinata) TERT gene was identified using the Basic Local Alignment Search Tool (BLAST) with the purple sea urchin TERT protein sequence as query against the *O. echinata* transcriptome contigs. The DNAWorks program was used to design 96 oligonucleotides to span the open reading frame of the brittle star TERT gene (Hoover and Lubkowski 2002). The synthetic gene was constructed by “oligo shuffling” as previously described with minor modifications (Stemmer et al. 1995). Oligonucleotides were pooled into four groups of 24 (4 µM each) and 1 µl used in a 25 µl PCR reaction with 1× Q5 Reaction buffer (25 mM TAPS-HCl at pH 9.3, 50 mM KCl, 2 mM MgCl₂, and 1 mM β-mercaptoethanol), 0.2 mM each dNTP, and 0.5 U of Q5 DNA Polymerase (NEB). One microliter of the previous PCR reaction was then used in a second 25 µl PCR reaction with 0.5 µM outermost oligonucleotides as primers. These PCR products were cloned into the pCITE4a vector with an N-terminal Flag epitope tag generating pN-Flag-oecTERT.

Telomerase in vitro reconstitution

The recombinant sea urchin and brittle star TERT proteins were synthesized from pN-Flag-spuTERT and pN-Flag-oecTERT, respectively, in a 5 µl reaction at 30°C for 60 min using the TnT Quick-coupled transcription/translation kit (Promega), following the manufacturer’s instructions. Various purple sea urchin and brittle star TR fragments were in vitro transcribed, gel purified, and added at a final concentration of 1 µM to assemble with the TERT protein in RRL at 30°C for 30 min.

Telomerase activity assay

Telomerase activity was assayed by the direct primer-extension assay (Li et al. 2013). A 10 µl reaction was performed with 2 µl in vitro reconstituted telomerase in 1× telomerase reaction buffer (50 mM Tris-HCl at pH 8.3, 2 mM DTT, 0.5 mM MgCl₂, and 1 mM spermidine), 100 µM dGTP, 100 µM dATP, 2 µM dGTP, 0.33 µM [α-32P]-dGTP (3000 Ci/mmole, 10 mCi/mL, Perkin-Elmer) and 1 µM (TTAGGG)₃ DNA primer. The reaction was incubated at 30°C for 1 h and terminated by phenol/chloroform extraction followed by ethanol precipitation. The telomerase extended products were resolved on a 10% polyacrylamide/8 M urea denaturing gel, and the dried gel was exposed to a phosphorstorage screen and analyzed with a Molecular Imager FX-Pro (Bio-Rad).

DATA DEPOSITION

The complete sequences of echinoderm TRs determined in this study have been deposited in the GenBank under accession numbers KT279799–KT279804. The echinoderm TR sequences identified from assembly of transcriptome data have been deposited in the Telomerase Database (Podlevsky et al. 2008).

SUPPLEMENTAL MATERIAL

Supplemental material is available for this article.

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