The teneurin C-terminal domain possesses nuclease activity and is apoptogenic.
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ABSTRACT
Teneurins are type 2 transmembrane proteins expressed by developing neurons during periods of synaptogenesis and apoptosis. Neurons expressing teneurin-1 synapse with other teneurin-1-expressing neurons, and neurons expressing teneurin-2 synapse with other teneurin-2-expressing neurons. Knockdowns and mutations of teneurins lead to abnormal neuronal connections, but the mechanisms underlying teneurin action remain unknown. Teneurins appear to have evolved via horizontal gene transfer from prokaryotic proteins involved in bacterial self-recognition. The bacterial teneurin-like proteins contain a cytotoxic C-terminal domain that is encapsulated in a tyrosine-aspartic acid repeat barrel. Teneurins are likely to be organized in the same way, but it is unclear if the C-terminal domains of teneurins have cytotoxic properties. Here we show that expression of teneurin C-terminal domains or the addition of purified teneurin C-terminal domains leads to an increase in apoptosis in vitro. The C-terminal domains of teneurins are most similar to bacterial nucleases, and purified C-terminal domains of teneurins linearize pCDNA3 and hydrolyze mitochondrial DNA. We hypothesize that yet to be identified stimuli lead to the release of the encapsulated teneurin C-terminal domain into the intersynaptic region, resulting in programmed cell death or the disruption of mitochondrial DNA and the subsequent pruning of inappropriate contacts.

KEY WORDS: Teneurin, Odz, Development, Apoptosis, Endonuclease, ABC complex

INTRODUCTION
Teneurins are type 2 transmembrane proteins, first discovered in Drosophila (Baumgartner and Chiquet-Ehrismann, 1993; Levine et al., 1994), that are expressed in the nervous system of vertebrates during periods of synaptogenesis and programmed cell death (Tucker and Chiquet-Ehrismann, 2006; Young and Leamey, 2009; Leamey and Sawatari, 2014; Mosca, 2015). The extracellular region of teneurins has a domain organization that is highly conserved across bilateria, with eight epidermal growth factor (EGF)-like repeats (Minet and Chiquet-Ehrismann, 2000), a rearrangement hot spot (RHS)-associated core protein-like domain with a predicted furin cleavage site (Tucker et al., 2012), and a C-terminal domain (CTD) like proteins contain a cytotoxic C-terminal domain that is encapsulated in a tyrosine-aspartic acid repeat barrel. Teneurins are likely to be organized in the same way, but it is unclear if the C-terminal domains of teneurins have cytotoxic properties. Here we show that expression of teneurin C-terminal domains or the addition of purified teneurin C-terminal domains leads to an increase in apoptosis in vitro. The C-terminal domains of teneurins are most similar to bacterial nucleases, and purified C-terminal domains of teneurins linearize pCDNA3 and hydrolyze mitochondrial DNA. We hypothesize that yet to be identified stimuli lead to the release of the encapsulated teneurin C-terminal domain into the intersynaptic region, resulting in programmed cell death or the disruption of mitochondrial DNA and the subsequent pruning of inappropriate contacts.

Phylogenetic analysis reveals that teneurins likely evolved through horizontal gene transfer from a bacterium into an ancestral choanoflagellate, with the eukaryotic proteins being formed through the fusion of a bacterial gene encoding a cysteine-rich domain, a six-bladed β-propeller, YD repeats, an RHS core-associated domain and a CTD with an eukaryotic gene encoding a type-2 transmembrane protein with EGF-like repeats (Tucker et al., 2012). Recent advances in our understanding of the structure and function of the teneurin-like prokaryotic proteins have generated significant insights into the structure and possible function of teneurins themselves. The prokaryotic proteins with YD repeats tend to be either type-2 transmembrane proteins that play a role in self-recognition between bacteria (Zhang et al., 2012; Koskinemi et al., 2013) or part of the ABC complex of secreted toxins that contribute to eukaryotic pathogenicity (Zhang et al., 2012; Hachani et al., 2014). Some transmembrane prokaryotic YD proteins are remarkably similar to teneurins in their domain architecture, though other domains substitute for the EGF-like repeats seen in teneurins (Fig. 1A). It has been hypothesized that identical YD proteins interact through their six-bladed β-propellers the C-terminal toxin domain is not released, but the toxin domain is released following heterotypic interactions. The toxins found in the CTD are diverse and include deaminases, nucleases and peptidases (Zhang et al., 2012). The secreted BC complex of the ABC toxin in the bacterium Yersinia entomophaga is homologous in organization to the extracellular region of metazoan teneurins from the six-bladed β-propeller distal to the CTD (Fig. 1A). Elegant structural studies revealed that the YD repeats of this toxin form a hollow barrel, with the β-propeller exposed at one end and the RHS repeat associated core
domain and toxic CTD stuffed into the other end (Busby et al., 2013) (Fig. 1B). The RHS domain of the BC proteins is cleaved as the protein folds, leaving the toxin free to diffuse away from a disrupted YD barrel (Busby et al., 2013). Thus, the cytotoxic CTD is encapsulated, protecting the bacterium and allowing for yet-to-be determined structural changes to cause the timely release of the toxin, either into the extracellular space or after uptake of the YD barrel into a target eukaryotic cell.

Is the CTD of teneurins, like the CTD of teneurin-like YD proteins, cytotoxic? Here we report the effects of the CTD of teneurin-1 and teneurin-2 on cell survival and further characterize the nature of its toxicity. Our observations lead us to predict that teneurins may act in the developing brain in a manner similar to their prokaryotic precursors.

RESULTS

The teneurin C-terminal domain is cytotoxic

To determine if the teneurin CTD is cytotoxic, HEK 293 cells were transfected with a plasmid expressing EGFP-linked human teneurin C-terminal sequences. The expressed sequences correspond to most of the RHS repeat-associated core-like domain including the predicted furin cleavage site and the entire CTD. The expressed protein corresponds to the part of the teneurin that would be released from a disrupted YD barrel if the protein is processed at the predicted furin cleavage site (for details see Materials and Methods and Fig. S1). For clarity, the expressed proteins are referred to below as ‘CTD’. The cells were allowed to grow for different periods up to 48 h, then fixed and stained with Crystal Violet. The density of the staining of control and experimental cultures is similar for the first 7 h, but fewer cells appear to be present in the cultures expressing EGFP-linked CTDs at later time points (Fig. 2A). Next we determined if the apparent loss of cells was due to apoptosis by immunostaining the cultures with an antibody to the cleaved form of caspase-3 (Fig. 2B). At 7 h this antibody recognizes a band of the appropriate molecular mass on immunoblots of lysates of CTD expressing cells, but not control cells (Fig. 2C). Cell counts show a statistically significant increase in the number of anti-cleaved caspase-3-expressing cells in transfected cells after both 48 h and 72 h; similar observations are made in COS-7 cells transfected with an expression construct designed to generate teneurin CTD sequences fused with a myc tag instead of the EGFP tag (Fig. 2D). Note that controls used in these experiments were cells expressing the empty vectors and not cells expressing an unrelated protein of similar size to the CTDs.

Next we determined if purified chicken teneurin CTDs are cytotoxic when added to tissue culture medium. The CTD from teneurin-1 at final concentrations as low as 0.05 µg/ml (∼4 nM), cause a significant increase in the proportion of cells that are positive for anti-cleaved caspase-3 after just 7 h, but higher concentrations are needed to see significant effects with the CTD of teneurin-2 (Fig. 2E). cDNAs made from RNA isolated from sorted HEK 293 cells expressing EGFP-tagged CTDs from teneurins-2 and control cells were used for transcriptional profiling (Table S1). Transcripts that are significantly up- or down-regulated include a DNA repair enzyme (RFC1) and factors that others identified in neurons induced to undergo apoptosis (ATF3, HDHD3) (Kristiansen et al., 2011; Chen et al., 2014). These and other changes were confirmed with RT-qPCR (Fig. 3).

Phylogenetic analysis of the teneurin C-terminal domain

Alignment of the sequences reveals the degree of identity and similarity between the GHH toxins of representative bacterial proteins
and the teneurin CTDs (Fig. 4). For example, the amino acid sequence of the GHH toxin of a teneurin-like YD repeat protein from Brevibacillus brevis and the CTD of human teneurin-1 are 23.37% identical and 28.57% similar, and the amino acid sequence of the GHH toxin domain of a non-YD repeat protein with a GHH toxin domain from Bacillus thuringiensis and the CTD of human teneurin-2 is 37.66% identical and 48.05% similar. Among tetrapods the CTD amino acid sequence is strongly conserved. For example, the 77 amino acid-long portion of the CTDs from human and chicken teneurin-1 that align with the GHH toxins share all but three amino acids. Matrices showing the identity and similarity of these domains are found in the supplemental information (Fig. S2).

The teneurin C-terminal domain has nuclease activity

Bacterial GHH toxins belong to the HNH/Endo VII family of endonucleases. Unlike the commonly encountered Type I and Type II restriction endonucleases that require Mg++ to catalyze the enzymatic activity, HNH endonucleases typically cleave DNA in the presence of either Ca++ or Mg++. As the GHH (or similar) amino acid motif, which is critical for the endonuclease activity of the bacterial toxins, has evolved into GYY or GYF in most vertebrate teneurins (Fig. 4), others have predicted that the teneurin CTD does not have endonuclease activity. We chose to test this by incubating the purified chicken C-terminal GHH toxin-like domains of teneurin-1 and teneurin-2 with the plasmid pcDNA3, a simple experiment based on the methods used by others (de Roodt et al., 2003), to demonstrate that certain components in snake venom have endonuclease activity. After 30 min the plasmid is cleaved by the purified teneurin-1 CTD, resulting in the appearance of linearized DNA together with open coil DNA (Fig. 5A). The GHH toxin-like CTD of teneurin-2 does not cleave the plasmid after 2 h, but does after an overnight incubation (Fig. 5A). If the plasmid is first linearized with EcoRI and then incubated with the teneurin-2 CTD, the CTD fails to further cleave the linear DNA (Fig. 5B). Thus, in short term experiments the GHH toxin-like CTD of teneurin-2 fails to demonstrate either endonuclease

Fig. 2. The teneurin-1 and teneurin-2 C-terminal domains are apoptogenic. (A) HEK 293 cells transfected with pEGFP alone or pEGFP with the CTD from human teneurin-1 (pEGFP-hTen1 CTD) or human teneurin-2 (pEGFP-hTen2 CTD) stained with Crystal Violet at different times following transfection. The control cultures become more densely stained over time, but not the cultures with cells expressing the CTDs. (B) Some of the cells transfected with the CTDs of teneurin-1 and teneurin-2 are immunostained with anti-cleaved caspase-3, a marker of apoptosis (arrows). (C) The anti-cleaved caspase-3 recognizes an appropriately sized band on an immunoblot of lysates from cells 7 h after being transfected with the teneurin-2 CTD (arrow). (D) A significantly higher percentage of HEK 293 cells expressing the EGFP-tagged CTDs are immunostained with anti-cleaved caspase-3 than controls at both 48 h and 72 h (n=3). Similarly, a higher percentage of COS-7 cells expressing myc-tagged CTDs are cleaved caspase-3-positive at these time points than cells transfected with pCMV6-A-puro alone (n=3). (E) The purified CTDs of both chicken teneurin-1 and chicken teneurin-2 cause a significant increase in the number of COS-7 cells that are positive for anti-cleaved caspase-3 when added to the culture medium (n=3). Error bars indicate mean±s.d. *P<0.05, **P<0.01, compared with controls (Student’s t-test).
or exonuclease activity, but it does exhibit endonuclease activity after longer incubation times. Controls included incubating the plasmid with an irrelevant protein of similar size that was purified using the same methods as the teneurin CTDs and incubating the teneurin-1 CTD with the plasmid in the presence of 10 mM EDTA. The irrelevant protein fails to cut the plasmid, and 10 mM EDTA, which should chelate the Mg++ needed for potentially contaminating Type I and Type II restriction endonucleases, does not inhibit the endonuclease activity of the CTD (Fig. 5C). This is consistent with the proposed relationship of the teneurin CTD to Ca++-dependent, and Mg++-independent, HNH/Endo VII family endonucleases. The sequence in pcDNA3 that is cleaved by the teneurin-1 CTD was determined by cloning linearized plasmid into the EcoRV site of pBluescript and sequencing with a T3 primer. The plasmid sequence that is cleaved (TACGACTCACTA|TAGGGAGACCCA) does not correspond to the sequence of any published restriction endonuclease. When the teneurin-1 CTD is incubated with the plasmid for longer periods (Fig. 5C), or when higher concentrations

| Name     | Gene ID     | FC  | Information                          |
|----------|-------------|-----|--------------------------------------|
| ATF3     | NM_001040619| 0.63| Activating transcription factor 3    |
| KDELRL3  | NM_006555   | 1.51| KDEL endoplasmic reticulum retention | rec 3 |
| KIF27    | NM_017576   | 0.33| Kinesin family member 27            |
| HDHD3    | NM_031219   | 1.55| Halolacid dehalogenase-like hydrolase|
| DNAH17   | NM_173628   | 0.32| Dynin heavy chain 17                |
| TUBB2C   | NM_006088   | 1.57| Beta-tubulin 4B                     |
| RFC1     | L23320      | 0.43| Replication factor C subunit 1      |

Fig. 3. Transcriptional profiling of HEK 293 cells expressing the C-terminal domain of teneurin-2 and confirmation by qRT-PCR. (A) The genes that were down-regulated in HEK 293 cells expressing the CTD from teneurin-2 were analyzed using online tools from the Gene Ontology Consortium. Many of the genes were related to the cytoskeleton and vesicle transport, enzyme modulation, cell signaling and nucleic acid binding. (B) A table listing seven representative genes that were either up-regulated or down-regulated in the HEK 293 cells following expression of the C-terminal domain of teneurin-2 showing the fold change (FC), their gene identification number and their common name. A complete list of up- and down-regulated genes is included as supplemental information (Table S1). (C-I) The changes found by transcriptional profiling of the representative genes were confirmed with qRT-PCR. (J,K) qRT-PCR also confirmed the overexpression of the CTDs from teneurin-1 and teneurin-2 in the HEK 293 cells used in the analysis. *P<0.05.
of the teneurin-1 CTD are incubated with the plasmid (Fig. 5D), the CTD appears to have exonuclease activity as well. To see if the CTD can cleave plasmid if it is encapsulated in the YD barrel, the entire extracellular domain of chicken teneurin-2 was purified and incubated with plasmid for up to 17 h. No nuclease activity was observed (Fig. 5E).

The teneurin C-terminal domain cleaves mitochondrial DNA in vitro

As mitochondria are needed to develop and maintain synapses (Li et al., 2004; D’Amelio et al., 2012; Hermes et al., 2016), and mitochondria would provide a convenient prokaryotic DNA-like target for toxins released near or into the developing synaptic cleft (Jiang et al., 2012), CTDs from teneurin-1 and teneurin-2 were incubated with a plasmid containing the entire murine mitochondrial DNA (pAM1). The CTD of teneurin-1 hydrolyzes the plasmid after 2 h, and the same domain from teneurin-2 hydrolyzes the mitochondrial DNA-containing plasmid after incubation overnight (Fig. 5F). Thus, both teneurin-1 and teneurin-2 have C-terminal GHH toxin-like domains with the potential to degrade mitochondrial DNA and initiate either failure of synaptogenesis or apoptosis.

DISCUSSION

The motif RXRR is found in all vertebrate teneurin sequences examined to date just C-terminal to the RHS domain. This motif is predicted in silico to be cleaved by furin or a furin-like protease (Tucker et al., 2012), but there is meager experimental evidence supporting this prediction. Antibodies raised against the CTD of Caenorhabditis elegans ten-1 immunostained cell membranes in nematode embryos, but the antibodies did not label blots of embryo homogenates (Drabikowski et al., 2005). A polyclonal antibody raised against the extracellular domain of mouse teneurin-1, which may have included antibodies to the CTD, recognized both high molecular weight bands and a band running near the bottom of the gel on immunoblots of mouse brain homogenates (Oohashi et al., 1999). While the smaller band may represent processed CTD, future experiments need to address this issue before broader speculation about the role of the CTD in neuronal development can be made.
that peptides corresponding to the C-terminal 38-41 amino acids of teneurins can modulate anxiety-related behavior and dendritic spine density in the hippocampus of rats following introduction of the peptide into the right lateral ventricle (Tan et al., 2011). These teneurin C-terminal associated proteins (TCAPs) have very different properties from the CTDs described here. For example, instead of inducing apoptosis when added to the medium of N38 cells, TCAPs protect the cells from necrosis by alkaline pH (Trubiani et al., 2007). In full-length teneurins, the TCAP sequence is C-terminal to, and does not overlap with, the region that alignment with YD proteins indicates is the likely site of DNA binding and nuclease activity within the CTD. Thus, there is no reason to expect that TCAPs have either nuclease activity or are cytotoxic. More importantly, TCAPs probably do not act as a functional domain of full length teneurins, as they can be transcribed independently (Chand et al., 2013).

Bacteria evolved toxins encapsulated in a YD barrel with a six-bladed β-propeller, as part of the struggle for limited resources between different strains and species. Following the acquisition of a YD repeat-containing protein by an ancestral chaoanflagellate, the early metazoan genome was primed for conducting a similar sort of chemical warfare between neurons in the central nervous system. We believe that the results presented here demonstrate evidence of a heretofore unknown mechanism for regulating neuronal development.

**MATERIALS AND METHODS**

**Phylogenetic analysis**

Teneurin CTDs are identified as GHH toxin-like domains by Pfam (http://pfam.xfam.org). For example, in the case of human teneurin-1 (NP_001156750) the CTD corresponds to the 78 amino acids found between amino acids 2648 through 2725. The CTDs of human teneurin-1 and teneurin-2 (NP_001073897) were used in an NCBI tBLASTn search of prokaryotic sequences to identify the most similar domains in bacteria. The most similar YD repeat proteins were GHH toxin containing proteins from *Brevibacillus brevis* (BAH45440) and *Bacillus cytotoxicus* (CP000764; 2072317-2072078). The most similar sequence overall was a secreted GHH toxin without YD repeats from *Bacillus thuringiensis* (AJH07449). The GHH toxin domains from these bacterial sequences were used in subsequent analyses. The CTDs of teneurin-1 through –4 sequences from human (*Homo sapiens*; XP_011529539; NP_001073897; NP_001073946; XP_011543235), mouse (*Mus musculus*; NM_011855; NM_011856; NP_035987; BAA77399), chicken (*Gallus gallus*; NP_994248; NP_001185466; XP_015136353), the elephant shark (*Callorhinus milii*; XP_007893009; XP_007900206; XP_007894102; XP_0079009700) as well as tena and tenam from the fruit fly (*Drosophila melanogaster*; NP_511137; CAAS51678) were aligned using Muscle 3.7 at Phylogeny.fr. Percent amino acid identity and similarity was determined using the SIAS program (http://imed.med.ucm.es/Tools/sias.html) of the Immunomedicine Group of the Universidad Complutense Madrid using the default amino acid grouping, the Blosum62 substitution matrix, and the cost for creating the gap Po set at 10 and imed.med.ucm.es/Tools/sias.html) of the Immunomedicine Group of the Universidad Complutense Madrid using the default amino acid grouping, the Blosum62 substitution matrix, and the cost for creating the gap Po set at 10 and the 125 C-terminal amino acids of teneurin-2; Fig. S1) were cloned from human cerebellum cDNA (AMS Biotechnology, Abingdon, UK, SRR306844) with the following primers: Teneurin-1 CTD (forward: ATGTCACTGTGTCCCATGACTTC, reverse: AGGCAGTCTTTGGAATAGCTGCTTGTTCTGCATG)

Others have found that the C-terminal 38-41 amino acids of human teneurins have sequence identity (up to 12%) with certain corticotropin-releasing factor peptides (Lovejoy et al., 2009), and

**C-terminal domain expression vectors, transfection and cell culture**

All cloning was done using traditional methods and the following kits and reagents: KOD Hot Start DNA polymerase (Novagen, Darmstadt, Germany, 71086-3), PCR product purification kit (Qiagen, 28106), gel extraction kit (Qiagen, 28706), Rapid DNA Ligation kit (Roche, 1163579001), and High Capacity cDNA Reverse Transcription kit (Applied Biosystems, Foster City, USA, 4322171). Teneurin-1 and teneurin-2 CTDs (the 123 C-terminal amino acids of teneurin-1, and the 125 C-terminal amino acids of teneurin-2; Fig. S1) were cloned from human cerebellum cDNA (AMS Biotechnology, Abingdon, UK, SRR306844) with the following primers: Teneurin-1 CTD (forward: ATGTCACTGTGTCCCATGACTTC, reverse: AGGCAGTCTTTGGAATAGCTGCTTGTTCTGCATG).
Violet for 1 h, gently rinsed in distilled water and allowed to air dry. These was fixed in 4% paraformaldehyde in phosphate buffered saline (PBS) for human teneurin-1 or -2 CTDs (see above) using jetPEI transfection reagent. 2 µg of the empty pEGFP vector or pEGFP vectors with inserts encoding the dishes in DMEM and 10% FBS. On day 2, the cells were transfected with hATF3 AAGTAAACCCCTTTGGACTTGAG); DNAH17 ACATGCACCACATTCGTGA, reverse: GGATTGGCAGAAAGGCTAAA); RAB7A GTAATTAAGCGGGACCAGCA, reverse: AGGTCCTCAGCTTTCGG-

Crystal Violet staining
On day 1, 2.5×10^5 HEK 293 cells were plated onto 3.5 cm tissue culture dishes in DMEM and 10% FBS. On day 2, the cells were transfected with 2 µg of the empty pEGFP vector or pEGFP vectors with inserts encoding the human teneurin-1 or -2 CTDs (see above) using jetPEI transfection reagent. Immediately following transfection one dish from each experimental group was fixed in 4% paraformaldehyde in phosphate buffered saline (PBS) for 30 min, then washed three times in PBS, then stained with 0.1% Crystal Violet for 1 h, gently rinsed in distilled water and allowed to air dry. These dishes represented time point ‘0 h’. Cells in the other dishes were cultured for 7 h, 24 h or 48 h and then processed in the same manner.

Transcriptional profiling and quantitative RT-PCR
Next, 1.2×10^6 HEK 293 cells were seeded in 10 cm tissue culture dishes in DMEM supplemented with 10% FBS. Cells were transfected with 12 µg of expression vector with jetPEI. After 16 h the cells were trypsinized and centrifuged at 1100 rpm for 3 min. The medium was removed and the cells were resuspended in 500 µl of PBS with 2% FBS, filtered (CellTrics; Partec; Goerlitz, Germany, 04-004-2327) and the green fluorescent cells were sorted. Then, 10×10^4 sorted cells were collected and RNA was isolated. RNA profiling was conducted and the data analyzed as described previously (Asparuhova et al., 2015). Quantitative RT-PCR was used to confirm selected profiles using Platinum SYBR Green qPCR SuperMix-UDG (Thermo Fisher Scientific, LC2002). Similar loading and transfer of proteins were confirmed by staining the membranes with Amido Black. After a 1 h blocking step in 5% milk powder in Tris-buffered saline with 0.05% Tween-20, membranes were incubated overnight with a rabbit polyclonal antibody to cleaved caspase-3 (Asp175) diluted 1:1000 (Cell Signaling Technology, 9611S). After incubation for 1 h with anti-rabbit IgG coupled to horseradish peroxidase (MP Biomedicals, Burlingame, USA, 0855689), blots were detected using SuperSignal (Thermo Fisher Scientific, 34080) followed by exposure to Kodak BioMax MR Films (Sigma-Aldrich, Z50397).

Immunoblotting and analysis of C-terminal domain induced apoptosis
Transfected HEK 293 cells were lysed in 4× sample loading buffer and separated by SDS-gel electrophoresis on a 12.5% polyacrylamide gel and electroblotted to polyvinylidene difluoride membranes (Thermo Fisher Scientific, LC1900). The membranes were developed using SuperSignal (Thermo Fisher Scientific, 34080) followed by exposure to Kodak BioMax MR Films (Sigma-Aldrich, Z50397). For protein expression and purification, the CTDs of chicken teneurin-1 and -2 were cloned using previously described eukaryotic expression protocols (Kozak, 2013). The cloning and expression vectors were transfected into HEK 293 cells in DMEM with 10% FBS and transfected as described above. Alternatively, COS-7 cells were seeded and purified CTDs from teneurin-1 or teneurin-2 were added to the medium at various concentrations. Cell cultures were fixed at various times following transfection or the addition of purified proteins with 4% paraformaldehyde in PBS for 30 min, washed twice in PBS, permeabilized for 5 min with 0.1% Triton in PBS, and rinsed again twice in PBS. Diluted anti-cleaved caspase-3 (see above) was added for 1.5 h at room temperature and the samples were then rinsed four times with PBS. For cultures expressing myc-tagged proteins, cultures 200 cells were incubated with mouse anti-myc for 1.5 h and then rinsed. The cells were then incubated in goat anti-mouse Alexa Fluor 488 or goat anti-rabbit Alexa Fluor 568 secondary antibodies (Molecular Probes) for 1 h. Studies with secondary antibodies alone were done in parallel to control for background. After final rinses and staining with Hoechst 33258 the cultures were mounted with ProLong Gold antifade reagent (Molecular Probes, 736934) and examined and photographed with an Axioskop photomicroscope (Carl Zeiss, Oberkochen, Germany) fitted with an ORCA-ER digital camera (Hamamatsu, Hamamatsu City, Japan). To analyze apoptosis in the transfected cultures, 200 cells from each experimental condition were identified with the nuclear stain and then scored as either untransfected and positive or negative for anti-cleaved caspase-3, or transfected (EGFP or anti-myc positive) and positive or negative for anti-cleaved caspase-3. To analyze apoptosis in the cultures with the purified proteins added to the medium, 200 cells were visualized in the medium and scored as positive or negative for the anti-cleaved caspase-3. Results from three independent experiments were analyzed using a Student’s t-test (one-tailed, unequal, paired variance).
**Endonuclease activity**

We incubated 200 ng of pcDNA3 (Thermo Fisher Scientific) for 2 h with various concentrations of the teneurin-1 CTD in the presence of 1× EcoRI buffer (New England BioLabs, Ipswich, USA, B0101S) to determine the optimal conditions for further studies (Fig. 5D). Further reactions included the pcDNA3 plasmid mixed with 500 ng of the CTDs of chicken teneurins-1 and -2 at 37°C for different periods of time, predigestion of the plasmid with EcoRI (stopping the reaction by heating to 65°C for 20 min prior to the addition of the purified teneurin domains), and including 10 μM EDTA in the reaction mix. For an additional control, pcDNA3 was incubated with a similarly sized protein corresponding to the 9th and 8th fibronectin type III domains of human tenasin-W, which was expressed and purified using the same conditions as the CTDs (Degen et al., 2007), and with the entire extracellular domain of chicken teneurin-2 (Beckmann et al., 2013). The sequence within pcDNA3 that was cut by the toxin domain of teneurin-1 was determined by gel purification of the linearized plasmid followed by blunt cloning into the EcoRV site of pBluescript (Stratagene, San Diego, USA), followed by sequencing using a T3 primer. The plasmid containing the complete DNA of murine mitochondria, pAM1 (Martens and Clayton, 1979), was used for some reactions.

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**Competing interests**

The authors declare no competing or financial interests.

**Author contributions**

Conceptionalization: R.P.T., R.C.E.; Methodology: J.F., R.P.T., R.C.E.; Validation: R.P.T.; Formal analysis: R.P.T.; Investigation: J.F., R.P.T.; Resources: R.C.E.; Writing - original draft: R.P.T.; Writing - review & editing: J.F.; Supervision: R.P.T., R.C.E.; Project administration: R.C.E.

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**Supplementary information**

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