A thrombospondin in the anthozoan *Nematostella vectensis* is associated with the nervous system and upregulated during regeneration

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Summary

Thrombospondins are multimeric extracellular matrix glycoproteins that play important roles in development, synaptogenesis and wound healing in mammals. We previously identified four putative thrombospondins in the genome of the starlet sea anemone *Nematostella vectensis*. This study presents the first analysis of these thrombospondins, with the goals of understanding fundamental roles of thrombospondins in the Eumetazoa. Reverse transcriptase PCR showed that each of the *N. vectensis* thrombospondins (Nv85341, Nv22035, Nv168100 and Nv30790) is transcribed. Three of the four thrombospondins include an RGD or KGD motif in their thrombospondin type 3 repeats at sites equivalent to mammalian thrombospondins, suggesting ancient roles as RGD integrin ligands. Phylogenetic analysis based on the C-terminal regions demonstrated a high level of sequence diversity between *N. vectensis* thrombospondins. A full-length cDNA sequence was obtained for Nv168100 (NvTSP168100), which has an unusual domain organization. Immunohistochemistry with an antibody to NvTSP168100 revealed labeling of neuron-like cells in the mesoglea of the retractor muscles and the pharynx. *In situ* hybridization and quantitative PCR showed that NvTSP168100 is upregulated during regeneration. Immunohistochemistry of the area of regeneration identified strong immunostaining of the glyocalyx, the carbohydrate-rich matrix coating the epidermis, and electron microscopy identified changes in glyocalyx organization during regeneration. Thus, *N. vectensis* thrombospondins share structural features with thrombospondins from mammals and may have roles in the nervous system and in matrix reorganization during regeneration.

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Introduction

Thrombospondins are multidomain, calcium-binding extracellular matrix (ECM) glycoproteins that have been most intensively studied in mammals, where they form a gene family of five members (TSP1–5). Mammalian thrombospondins have important roles in angiogenesis, wound healing, tumor progression, cardiovascular biology, the development and maintenance of connective tissues and synaptogenesis. These functions are mediated by interactions with many binding partners including other ECM components, growth factors, matrix proteases, integrins, proteoglycans and other adhesion and signaling receptors (for reviews see Adams and Lawler, 2012; Risher and Ergul, 2012). Along with SPARC, thrombospondins are the most ancient modulatory adhesion molecules, with many interactions that relate to ECM assembly and remodeling (Mosher and Adams, 2012). In structure, the thrombospondins of mammals fall into two subgroups, A and B, that differ in domain composition in their N-terminal regions and assemble through the action of coiled-coil domains either as homo-trimers (subgroup A; TSP1 and TSP2) or as homo-pentamers (subgroup B; TSP3, TSP4 and TSP5). Gene conservation and similarities of expression patterns suggest that the TSP1–5 family has related functions in many other cnidarians (Tucker, 1993; Tucker et al., 1995; Urry et al., 1998; McKenzie et al., 2006; Kasprick et al., 2011).

Although many important ECM components are specific to gnathostomes (jawed vertebrates) (Özbek et al., 2010), it is now appreciated that thrombospondins are conserved through most metazoan phyla. The genomes of the purple sea urchin *Strongylocentrotus purpuratus* and the sea squirt *Ciona intestinalis* contain thrombospondin genes encoding proteins with subgroup A and B domain architectures as well as a protein belonging to an additional subgroup termed DD due to the presence of a “discoidin-like domain” at the N-terminus (Bentley and Adams, 2010). In contrast, most protostomes and early-diverging metazoan species encode a single thrombospondin with, in general, subgroup B-like domain architecture (Bentley and Adams, 2010). Functional studies of thrombospondins in invertebrates have only been carried out in *Drosophila melanogaster*. *D. melanogaster* thrombospondin is a pentameric protein that undergoes interactions with heparin and PS2 integrin...
and is functionally essential for the development of muscle–tendon attachment sites (Adams et al., 2003; Chanana et al., 2007; Subramanian et al., 2007; Gilsohn and Volk, 2010). These studies indicate conservation of ECM-related functions and interactions of thrombospondins within bilaterians, yet to date nothing is known about the thrombospondins of diploblasts or other early-diverging metazoa. In view of the high conservation of thrombospondins across the metazoa, such investigations are important for understanding the fundamental roles of thrombospondins in tissues.

Cnidaria form a sister group to bilaterians and are increasingly recognized as important model organisms for studying the genomic and organismal evolution of animals. In contrast to D. melanogaster and Caenorhabditis elegans, which have undergone extensive gene losses, sequencing of several Cnidarian genomes has illuminated relationships with vertebrates and the possible complexity of the metazoan ancestor (Putnam et al., 2007; Chapman et al., 2010). Cnidaria are defined by having two epithelial cell layers (i.e. they are diploblastic), a single oral body opening and tentacles, a muscular system and nerve net, and cnidoblasts. The outer epidermis and inner gastrodermis are separated by an ECM layer, the mesoglea, that has features of both connective tissue ECM and basement membranes (e.g. Schmid et al., 1999; Tucker et al., 2011). The anthozoan Cnidarian Nematostella vectensis is an ustearine sea anemone that is emerging as a tractable laboratory model with a simple lifecycle (Hand and Uhlinger, 1992; Darling et al., 2005). As part of an overview of the evolution of metazoan ECM, we previously identified four thrombospondin genes in N. vectensis (Bentley and Adams, 2010; Özbek et al., 2010). Here we present the first analysis of these thrombospondins and their phylogenetic relationships. We report that a Nematostella thrombospondin with an unusual domain composition is expressed in adult polyps and is sharply upregulated during regeneration. The possible functions of this anthozoan thrombospondin that can be surmised from its distinctive patterns of expression indicate interesting parallels and differences with the thrombospondins of vertebrates.

Results

The four thrombospondins of Nematostella vectensis

BLAST searches of the N. vectensis genome identified four open reading frames that include regions characteristic of thrombospondins, i.e. EGF-like domains, thrombospondin type 3 repeats and a C-terminal L-lectin-like domain. These thrombospondins were previously referred to by us using the JGI transcript identification number preceded by “Nv” (Bentley and Adams, 2010; Özbek et al., 2010), and we continue to use that nomenclature here. Nv85341 (which corresponds to GenBank XP_1639928) is encoded on genome scaffold 11 and Nv22035 is on scaffold 62 (scaffold numbers refer to the JGI N. vectensis genome assembly) (Putnam et al., 2007). Nv168100 (which corresponds to GenBank XP_1631622) and Nv30790 are both encoded on scaffold 99, suggesting that these genes may be the result of a lineage-specific gene duplication. Inspection of the domain organizations of all four predicted proteins indicates that none of the predictions corresponded to a full-length thrombospondin sequence (Fig. 1A). The prediction for Nv30790 was extended at the N-terminal end through an overlapping expressed sequence tag (EST), FC258725. The region encoded by the EST included two cysteine residues in a position typical of the pentamerizing thrombospondins (Fig. 1A) (Bentley and Adams, 2010). We confirmed that all four genes are transcribed in adult N. vectensis using reverse transcriptase PCR (RT-PCR) (Fig. 1B).

Nematostella vectensis thrombospondins have RGD motifs

The tripeptide RGD acts as an integrin binding motif in some thrombospondins and many other ECM molecules (Ruoslahti, 1996; Bentley and Adams, 2010). RGD and KGD motifs are found in the thrombospondin type 3 repeats of three of the four predicted N. vectensis thrombospondins (asterisks in Fig. 1A); these sequences were confirmed in the sequenced RT-PCR products described above. Alignment with the thrombospondin type 3 repeat sequences of human thrombospondins (Fig. 2) demonstrated that the RGD and KGD motifs found in N. vectensis thrombospondins are in phylogenetically conserved locations. For example, RGD motifs are found in the second thrombospondin type 3 repeat of human TSP4 and 5, and an RGD motif is found at the same site in the second thrombospondin type 3 repeat of Nv30790. A KGD motif is found in the sixth thrombospondin type 3 repeat of human TSP1 that aligns with an RGD motif in Nv168100 and a KGD motif in Nv22035. Finally, RGD motifs are found in the seventh thrombospondin type 3 repeats of human TSP1 and TSP2 and these align with an RGD motif in Nv22035. It is interesting to note that the 14 residue RGD-containing sequence DTGDGRGDACDDD found in the sixth thrombospondin type 3 repeat of Nv168100: 1) is identical to the 14 amino acid (aa) RGD-containing sequence found in the second thrombospondin type 3 repeat of human TSP4, 2) is 93% identical to this same region in human TSP5 (i.e. 13 of the 14 residues are identical), and 3) contains the same string of nearby aspartic acids.

Molecular cloning and analysis of a Nematostella vectensis thrombospondin

The predicted Nv168100 appeared to be relatively complete, so it was selected for further analysis. The 1151 aa full-length coding sequence (NvTSP168100; GenBank accession number JX680803) (Fig. 1C) was obtained from overlapping PCR products generated using primers based on predicted open reading frame sequences as well as potential open reading frame sequences 3’ and 5’ to the sequences encoding the predicted protein. Following the signal peptide (aa 1–21), the N-terminal domain (aa 22–227) corresponds to a laminin G-like domain, which is canonical for the majority of thrombospondins (Tan et al., 2006; Bentley and Adams, 2010). The sequence from aa 228 to aa 471 is unique and includes a region from approximately aa 228 to aa 329 that is rich in proline, arginine and threonine residues (designated “PRT region”) and a region from aa 330 to aa 471 that includes a short predicted coiled coil. BLAST searches of GenBank with the PRT region identified proline-rich regions of around 50% identity in C. briggsae XP_002636096 (Fig. 1D) and multiple other hypothetical proteins of C. briggsae, and proline-rich regions of around 29–32% identity in a bacterial zinc metalloproteinase (ZP_04776628) and a bacterial cell wall anchor domain protein (ZP_06282692). The functions of these related polypeptide regions are unknown. The C-terminal region of NvTSP168100 consists of four EGF-like domains, a series of thrombospondin type 3 repeats and an L-type-lectin-like domain and thus
corresponds to a canonical thrombospondin C-terminal region (Fig. 1A,C).

**Phylogenetic analyses**

Thrombospondins are widespread throughout the metazoa and *N. vectensis* is unusual as an early-diverging metazoan that encodes multiple thrombospondins (Bentley and Adams, 2010). Phylogenetic analyses were carried out to examine the relationships of *N. vectensis* thrombospondins to previously identified thrombospondins. Sequences corresponding to the conserved C-terminal regions (comprising three EGF-like domains, the type 3 repeat, and the L-lectin domain) (Fig. 1A) were aligned by the MUSCLE algorithm and analyzed for phylogenetic relationships by neighborhood-joining or maximum-likelihood methods. *Nv22035* was excluded from this analysis because the sequence prediction does not include the complete C-terminal region (Fig. 1A). The tree based on neighborhood-joining analysis was rooted between the TSP-DD group, which in these analyses also included the thrombospondin from the placozoan *Trichoplax adhaerens* and *Nv85341*, and all other thrombospondins (Fig. 3A). This tree demonstrated a close relationship between *Nv116800* and *Nv30790*, consistent with the proposed origin from a lineage-specific gene duplication. The tree also indicated high diversity of the *N. vectensis* thrombospondins in that *Nv116800* and *Nv30790* appeared remote from *Nv85341* and other thrombospondins (Fig. 3A). Maximum-likelihood analysis with boot-strapping extended these indications. The unrooted tree (Fig. 3B) confirmed the separate and robust groupings of A and DD group thrombospondins and the grouping of *T. adhaerens* thrombospondin and *Nv85341* with the DD group. In this analysis, *Nv116800* and *Nv30790* clustered with the subgroup B thrombospondins of bilaterian protostomes and deuterostomes to form a large B group, albeit the rooting of this grouping was not well-supported by boot-strapping. Both methods indicated consistently that the C-terminal region of the thrombospondin from the demosponge *Amphimedon queenslandica* is remote from other thrombospondins; the maximum-likelihood analysis placed AqTSP on a separate branch from all other groups (Fig. 3B). To include *Nv22035* in the analyses, an unrooted tree was also prepared based on only
the type 3 repeats and L-lectin domain. This analysis placed Nv22035 separately from the previously described groups, not unlike the thrombospondin from *A. queenslandica*, and also placed the Nv116800/Nv30790 branch as a separate lineage related to both the A and B groups of TSPs (Fig. 3C).

The expression of a *Nematostella vectensis* thrombospondin

To localize NvTSP168100 in intact adult animals, a polyclonal antibody was raised in chickens against a portion of the N-terminal domain. The antibody was first characterized by immunoblotting. The polyclonal IgY recognized the antigen NvTSPN (Fig. 4A); this band was not seen following preabsorption of the antibody with antigen, nor was it present when the blot was incubated in preimmune IgY (not shown). The predicted molecular mass of NvTSP168100 is 130 kDa, and on immunoblots of the first pellet fraction (P1) of homogenized adult *N. vectensis* anti-NvTSP168100N recognized two bands running between 130 kDa and 250 kDa (arrowheads, Fig. 4A) and a third band running between 72 kDa and 95 kDa (arrow, Fig. 4A). These bands were not detected either with the preimmune IgY or after preabsorption of the antibody with antigen (not shown). The larger bands likely represent full length NvTSP168100 and are running higher than predicted due to posttranslational modifications such as glycosylation; the smaller band may result from proteolytic processing or may be a degradation product.

Immunohistochemistry was used to determine the sites of NvTSP168100 expression. Cross sections through the body column (scapus) of adult *N. vectensis* were incubated with anti-NvTSP168100 or with preimmune IgY and counterstained with TRITC-phalloidin (to show regions where myonemes, the cellular processes filled with retractile elements resembling striated muscle, are concentrated) and the nuclear stain DAPI. Immunostaining was seen in the mesoglea of retractor muscles in each of the mesenteries (i.e. the double folds of gastrodermis that project into the gastrovascular cavity) (Fig. 4B); the nearby parietal muscles were unlabeled as was the mesoglea found in adjacent sections incubated with preimmune serum (Fig. 4C). Adjacent sections incubated with preabsorbed anti-NvTSP168100N were also unlabeled (Fig. 4D). In longitudinal sections through the scapus of adult *N. vectensis* the nature of the immunolabeling of retractor muscles was clearer: the antibody labels long, beaded filaments or cell processes found between the longitudinally arrayed leaflets of the retractor muscle myonemes (Fig. 4E). In an oblique section through the mesoglea underlying the pharynx the anti-NvTSP168100N labeled cell bodies with DAPI-stained nuclei and long narrow processes embedded within the mesoglea. The morphology of these cells and their processes was examined further by confocal and transmission electron microscopy. Confocal observations (Fig. 4G) revealed neuron-like triangular cell bodies labeled with anti-NvTSP168100N that are embedded within the mesoglea. Long, narrow, beaded processes with occasional branches extend from the cell bodies. Thin sections through the mesoglea of adult polyps revealed numerous cellular processes running longitudinally within the mesoglea of the retractor muscles (Fig. 4H). These processes have a relatively uniform cross sectional diameter of 300–350 nm. As observed in the confocal microscope, the processes were often found in loose bundles, though sometimes they were found singly. Higher magnification showed that these processes are closely associated with ECM within the mesoglea (Fig. 4H, inset). Most of the processes contain large membrane-bound vesicles and numerous small (30–40 nm) dark granules. In some of the processes microtubules were seen in cross section closely associated with the large vesicles. Similar processes were not observed in the mesoglea of the parietal muscles or the body wall (Tucker et al., 2011). Thus the cells that were immunostained with anti-NvTSP168100N bear a strong morphological resemblance to neurons and have processes that contain elements characteristic of neurites. These neuron-like cells are limited to the mesoglea of the pharynx and retractor muscles, where they are closely associated with the myonemes.
The expression of a *Nematostella vectensis* thrombospondin is upregulated during regeneration.

We have shown previously that the ultrastructural appearance of the mesoglea in *N. vectensis* changes dramatically 48 hours after transection of the polyp (Tucker et al., 2011). To determine if the expression of NvTSP168100 changes during regeneration, 6-week-old polyps were transected and then processed for whole mount *in situ* hybridization 48 hours later. A signal was observed in the entire scapus, but the signal was strongest near the site of regeneration (Fig. 5A,B). To confirm upregulation of NvTSP168100 transcripts during regeneration we conducted quantitative PCR (qPCR) with NvTSP168100-specific primers and compared the level of expression in control polyps with the level of expression 48 hours after transection. qPCR confirmed the results of the *in situ* hybridization experiments, revealing a 41.8±11.1 (n=3) fold increase in the levels of NvTSP168100 mRNA following transection (Fig. 5C) (results were normalized to levels of a control rRNA). Next, immunohistochemistry was conducted on longitudinal sections through the scapus of polyps 48 hours following transection in order to confirm the results above and to localize the thrombospondin. The immunostaining of the neuron-like elements described above that are found within the mesoglea of the retractor muscles was still observed, but additional immunostaining was seen in the carbohydrate-rich lining of the epidermis known as the glycocalyx (Fig. 5D). This staining extended proximally toward the oral end of the column, where it abruptly ended (Fig. 5E). To determine if ultrastructural changes to the glycocalyx accompany regeneration, thin sections of the glycocalyx (preserved by using a fixative mixed with tannic acid and ruthenium red) were observed in the transmission electron microscope. In controls, the glycocalyx appeared as an electron-dense, fuzzy coat, 40–50 nm thick that lined the outer surface of the epidermis as well as the 1.5 μm long microvilli that protrude from the epidermis (Fig. 6A,B). This region was dramatically different 48 hours following transection: the microvilli were approximately twice as dense (3.2 microvilli/μm versus 1.4 microvilli/μm) and were often branched. Thin fibrils projected from the surface of the microvilli and often spanned the area between adjacent microvilli (Fig. 6C,D). Electron-dense debris appeared to be trapped at the outer limit of the microvilli and glycocalyx (Fig. 6B, arrows) and spherical elements ranging from 30–40 nm in diameter, which resembled viral particles, were associated with microvilli (Fig. 6D, arrowheads).

**Fig. 3.** Phylogenetic relationships of *N. vectensis* thrombospondins and other representative thrombospondins. In A and B, sequences of 620 residues including the last three EGF-like domains, thrombospondin type-3 repeats and the L-lectin-like domain from representative invertebrate and human thrombospondins were aligned in MUSCLE 3.7. (A) Rooted tree prepared by the neighborhood joining method in Phylip with proportionate branch lengths. (B) Unrooted tree prepared by the maximum-likelihood method in PHYML 3.0 with 200 cycles of bootstrapping. Bootstrap values are shown for each node: values above 0.7 are taken to indicate stability of the branchpoint. Scale bars show substitutions/site. *N. vectensis* thrombospondins are indicated by asterisks. (C) Unrooted tree prepared as in B, using only the sequences corresponding to the type 3 repeats and the L-lectin-like domain from representative invertebrate and human thrombospondins were aligned in MUSCLE 3.7. (A) Rooted tree prepared by the neighborhood joining method in Phylip with proportionate branch lengths. (B) Unrooted tree prepared by the maximum-likelihood method in PHYML 3.0 with 200 cycles of bootstrapping. Bootstrap values are shown for each node: values above 0.7 are taken to indicate stability of the branchpoint. Scale bars show substitutions/site. *N. vectensis* thrombospondins are indicated by asterisks. (C) Unrooted tree prepared as in B, using only the sequences corresponding to the type 3 repeats and the L-lectin-like domain so as to include Nv22035. Aq, *Amphimedon queenslandica*; Br, *Branchiostoma floridae*; Ci, *Ciona intestinalis*; Dp, *Daphnia pulex*; Dm, *Drosophila melanogaster*; Hs, *Homo sapiens*; Lg, *Lottia gigantea*; Nv, *Nematostella vectensis*; Sp, *Strongylocentrotus purpuratus*; Ta, *Trichoplax adhaerens*. 

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# A Nematostella thrombospondin

By guest on January 12, 2018
Discussion

Based on the fossil record and molecular data, the divergence of Cnidaria and the Bilateria is estimated to have occurred around 750 million years ago, with Cnidaria subsequently evolving as a single phylum and the bilaterian ancestor giving rise to over 30 phyla (Ryan et al., 2007). In view of the diversity of bilaterians, the genome sequence of *N. vectensis* has proved very informative in illuminating the genomic complexity of the Eumetazoan ancestor. In relation to the protostomes examined to date, the gene content of *N. vectensis* includes large sets within gene families such as the Hox genes (Ryan et al., 2007). The encoding gene content of *N. vectensis* also encodes unexpectedly large families of SPARC-like proteins (Koehler et al., 2009) and integrins (Knack Adams, 2010). Whether either Nv85341 or *T. adhaerens* thrombospondin contain an N-terminal discoidin domain must wait further sequencing studies, but it is interesting to consider the model that the DD group C-terminal region might have diverged as a separate lineage from the inferred metazoan ancestor (Fig. 3C). Nv85341, along with *T. adhaerens* thrombospondin, was most related to the DD group, previously identified only in early-diverging deuterostomes (Bentley and Adams, 2010). Whether either Nv85341 or *T. adhaerens* thrombospondin contain an N-terminal discoidin domain might have originated in the Eumetazoan ancestor, with subsequent loss from most protostome lineages. In line with other analyses of *A. queenslandica* gene content (Srivastava et al., 2010), the C-terminal region of *A. queenslandica* thrombospondin was distinct from all other thrombospondin sequences in the dataset. In the phylogenetic analysis of the shorter sequence region, including only the type 3 repeats and L-lectin domain, Nv22035 did not associate with any of the major thrombospondin groups and likely represents a lineage-specific divergence. This analysis, based on less sequence information, also moved Nv85341 to the edge of the DD group, but the central conclusion of great diversity of thrombospondin sequences within *N. vectensis* is upheld. Future analyses that include information from the yet-to-be sequenced genomes of other classes of cnidarians (e.g. the scyphozoans and cubozoans), as well as more complete *N. vectensis* sequences, may clarify these relationships.
lineage-specific forms distinct from the functional groups recognized to be conserved in Bilateria (Knack et al., 2008). Thus, it cannot be predicted which of these integrins might bind to *N. vectensis* thrombospondins. NvTSP168100 has a canonical laminin G domain-like N-terminal domain containing motifs rich in basic residues and can be predicted to bind heparan glycosaminoglycans.

Complete predicted polypeptide sequence was obtained for NvTSP168100. The general domain organization has some unique features that make it unlike group B thrombospondins in other taxa. Most notable is the PRT region found between the N-terminal domain and the predicted coiled coil. This sequence aligns well with sequences from predicted proteins of unknown function in the nematode *C. briggsae* (Fig. 1D), and to a lesser extent with cell-wall proteins from some prokaryotes, but these similarities do not shed light on its potential function at this time. However, proline–arginine rich peptides derived from a small leucine-rich proteoglycan family member (PRELP) have been identified to have antimicrobial properties (e.g. Malmsten et al., 2006), which is intriguing given that NvTSP168100 appears to be upregulated in the glycocalyx after wounding and during regeneration. NvTSP168100 contains only a short coiled coil and is also lacking the characteristic cysteine residues that form interchain disulfide bonds adjacent to the oligomerization region in the pentameric group B thrombospondins. In the future, the oligomerization state of NvTSP168100 would need to be identified by biochemical analysis. Nv30790, the predicted thrombospondin that is closely related to NvTSP168100 in its C-terminal region (Fig. 2), does contain the characteristically spaced cysteine residues that are predicted to form interchain disulfide bonds, thus this feature of subunit assembly is highly conserved, albeit not in all *N. vectensis* thrombospondins. The domain organization of NvTSP168100 suggests that the encoding gene underwent domain shuffling resulting in the inclusion of the PRT region after a lineage-specific gene duplication event that gave rise to Nv30790 and NvTSP168100.

An antibody raised against the N-terminal sequences of NvTSP168100 labeled neuron-like cells embedded in the mesogloea of retractor muscles and the pharynx in adult *N. vectensis*. Morphologically these cells have the hallmark features of neurons, especially their long, beaded processes (which are indistinguishable at the level of the light microscope from the neuronal processes of *Hydra* (Grimmelikhuijzen and Westfall, 1995)) and the presence of dense-cored vesicles and longitudinally arrayed microtubules within these processes. The labeled cells could not be identified conclusively as neurons by double labeling with neuron-specific probes, as such probes have not yet been characterized in adult *N. vectensis*. Further studies in the adult with antibodies to neurotransmitters such as RFamide that identify subsets of neurons in planula larvae and juvenile polyps (Marlow et al., 2009) are called for. Neuronal immunostaining for a thrombospondin is not entirely unexpected, as others have shown that anti-thrombospondins readily label neurons in the central nervous system of mammals (e.g. Arber and Caroni, 1995). What is surprising is the presence of a population of neuron-like cells within the mesoglea, as ultrastructural analysis of adult anthozoans typically describes neuronal perikarya within the epidermis or gastrodermis (reviewed by Grimmelikhuijzen and Westfall, 1995). The mesoglea of *N. vectensis* is distinct from that of many other Cnidaria in containing cells, but to date the major cell type...
recognized within the mesoglia is the amoebocyte, a migratory cell of the innate immune system (Frank and Bleakney, 1976; Tucker et al., 2011). Experiments with a promoter reporter transgene have suggested that the retractor and tentacles muscles have different gene expression profiles to other myoepithelial cells; these might include cues that promote nerve–muscle interactions in this body region where fast, dynamic movements are very important for prey engulfment (Renfer et al., 2010).

Very little is known about the molecular events that mediate regeneration in anthozoans. It is recognized that the process differs from wound repair (e.g. wounds that do not transect the body column do not lead to regeneration) and has features in common with events of embryogenesis. In N. vectensis, as in other Cnidaria, canonical Wnt signaling appears to have an important role in controlling regeneration (Treviso et al., 2011). ECM would be expected to have a role in regeneration and, indeed, in transected adult N. vectensis, morphological reorganization of the mesoglia is apparent in the region adjacent to the transaction site (Tucker et al., 2011). Here, we demonstrate that the epidermal glyocalyx is also reorganized during regeneration and that NvTSP168100 is a likely participant in this reorganization, as demonstrated by its upregulation and prominent altered localization. Further studies of the cellular and biochemical properties of NvTSP168100, as well as understanding of the other N. vectensis thrombospondins, will provide additional insights into the evolution of tissues and homeostatic cell–ECM interactions.

Materials and Methods

Nematostella culture

A colony of mature N. vectensis polyps descended from clones CH2 and CH6 (Hand and Uhlinger, 1992) were obtained from the University of California Davis Bodega Bay Marine Laboratory. The sea anemones were maintained in microfiltered sea water diluted to 33% with distilled water at room temperature and adjusted to pH 7.8. Details of raising the sea anemones can be found in Darling et al. and Tucker (Darling et al., 2005; Tucker, 2010), and details associated with transecting the polyps to study regeneration can be found in Tucker et al. (Tucker et al., 2011).

Gene identification, expression analysis and molecular cloning

Putative thrombospondin genes were identified in the genome of N. vectensis (Putnam et al., 2007) by BLAST (http://blast.ncbi.nlm.nih.gov) searches of JGI transcripts and by using the Superfamily database (http://superfamily.cs.bris.ac.uk). BLAST searches (SUPERFAMILY) searching for predicted proteins with multiple domains characteristic of thrombospondins that were present in the right order. Further domain analysis of the predicted proteins was carried out using Pfam (http://pfam.sanger.ac.uk) and SMART (http://smart.embl-heidelberg.de). The expression of the putative thrombospondins in adult N. vectensis polyps was confirmed using reverse transcriptase polymerase chain reaction (RT-PCR). Methods for polyA mRNA isolation and PCR amplification conditions were described previously (Tucker, 2010). The primers used to amplify products corresponding to the putative thrombospondin sequences identified above were: 5’-GATCAGCGGCTTACACAGAG-3’ and 5’-CGTGTCATCTTGTCGCTGT-3’ (Nv168100, XP_1631622), 500 bp product; these primers were also used as described below and called NvTSP168100F/R); 5’-CCGGTCTTTTATCCGGACATGG-3’ and 5’-TACCATACACCTGCCTATAC-3’ (Nv85341, XP_1639928; 499 bp product); 5’-GACTTGATGATGATCCAGA-3’ and 5’-CCG-TCGATGATGATCCAGA-3’ (Nv85340; 503 bp product); and 5’-TACAAC-3’ and 5’-TACCATACACCTGCCTATAC-3’ (Nv30790, 501 bp product). Each of the primer pairs above spanned introns to control for DNA contamination. In addition, a control primer pair corresponding to n-tubulin sequences was used (Tucker, 2010). Primers were ordered from Eurofins MWG Operon and size standards were purchased from Invitrogen. The PCR products were subcloned using a TOPO TA-Cloning kit (Invitrogen) into pCRII or pCR2.1 cloning vectors and sequenced by Davis Sequencing.

One of the putative thrombospondins (Nv168100, XP_1631622) was cloned to generate a potentially complete cDNA sequence using primers based on the predicted transcript as well as potential open reading frame sequences identified from the genome both 3’ and 5’ to the NvTSP168100F/R described predicted transcript to create five overlapping PCR products: NvTSP168100N2, NvTSP168100N, NvTSP168100 (the PCR product described above), NvTSP16800N and NvTSP168100C. This expanded the predicted protein to include a thrombospondin-like-N-terminal domain with a potential in-frame start codon and signal peptide and a complete L-lectin domain with an in-frame stop codon. The primer pairs used to generate overlapping PCR products were: NvTSP168100N2 (5’-CTGCTATGCGCTCACGG-3’), NvTSP168100N (5’-CTGATGCTTCTCTCCCTCTC-3’) and NvTSP16800NF (5’-AGTCCACACGGCGTCCATTTACCACT-3’), NvTSP168100N (5’-CCGCCTGCAATTTCTCAG-3’), NvTSP168100F (5’-GATACCACCCTTACACAGAG-3’), NvTSP168100 (5’-CCGTGACTTGTTGGCTGTG-3’), NvTSP16800CF (5’-TGATAACCTGCAACGGGAAAC-3’), NvTSP16800CR (5’-TCGTTGTCCTGCCAGAGATCCTAC-3’), NvTSP167100C (5’-TGGATGCAGTTGTTGTTGTTGTTG-3’). The PCR products were subcloned and sequenced as described above. The complete cDNA sequence is deposited as GenBank accession number JX680803.

Bioinformatics and phylogenetic analyses

Domain architectures of the predicted N. vectensis thrombospondin polypeptides were analyzed against the InterProScan database (Quevillon et al., 2005) at EBI (http://www.ebi.ac.uk/Tools/pfa/interproscan). Coiled-coil domains were identified with the COILs algorithm, with 2.5 fold weighting applied for the a and d positions (Lupas et al., 1991). Domain boundaries were assigned by MUSCLE (Edgar, 2004) multiple sequence alignment of the N. vectensis thrombospondins against well-characterized thrombospondin domains. Phylogenetic analyses were based on a previously developed dataset of thrombospondin sequences (Bentley and Adams, 2010). This dataset was curated for the current study by inclusion of Amphimendon queenslandica thrombospondin (GenBank XP_003389217) (Stivrastava et al., 2010) and limitation of vertebrate thrombospondin sequences to human TSP1–5. Multiple sequence alignment for the analysis was performed by BLAST (http://blast.ncbi.nlm.nih.gov) searches of JGI sequences was used (Tucker, 2010). Primers were ordered from Eurofins MWG Operon and size standards were purchased from Invitrogen. Protein expression and antibody production

Plasmid NvTSP168100N2 (pTSPN) was the source for DNA sequences used to express a recombinant partial N-terminal thrombospondin domain of N. vectensis for antibody production. pTSPN was digested with EcoRI to release an ~1.1 kb fragment encoding as 67 to aa 437 of the presumptive protein. This fragment was cloned into pT7-1 (pT7-7) ( Studier et al., 1990) with a one amino acid deletion to change the reading frame. The resulting plasmid was sequenced to verify DNA sequence and orientation then introduced into BL21-AI (Invitrogen) for large scale protein production. Protein expression was induced by the addition of 0.5 g of arabinose to 500 ml of exponentially growing E. coli at an OD600 ~0.8. After 5–6 hours of induction cells were harvested by centrifugation and stored as wet pellets at ~20°C until processed. The recombinant protein readily formed inclusion bodies that were purified using lysozyme/detergent lysis with DNase/RNase digestion followed by high salt and low salt washes (Nagai and Thogersen, 1987). Purified inclusion bodies were resuspended in 10 mM Tris pH 7.5, 1 mM EDTA, 0.2 M NaCl, 8 M urea and chromatographed on a GE Healthcare HiLoad 16/60 Superdex 200 gel filtration column. Fractions were monitored by SDS-PAGE and peak fractions pooled. Protein concentration was determined using a Pierce BCA assay with BSA as a standard (Thermo Scientific). Recombinant protein was dialyzed against PBS overnight to remove urea; the majority of the protein precipitated.

Purified chicken antibodies (made from 3 eggs prior to injection) and purified chicken anti-NvTSP168100N IgY were prepared at Aves Labs.

Paraffin embedding and immunohistochemistry

Adult N. vectensis polyps were immersed in 75% MgCl2 in 33% sea water for 20 minutes, fixed in 2% paraformaldehyde in phosphate buffered saline (PBS) at room temperature for 30 minutes, then fixed for 5 hours in 4% paraformaldehyde in PBS at 4°C. After rinsing overnight in cold PBS the specimens were trimmed using a razor blade and dehydrated in increasing concentrations of ethanol (50%, 75%, 95%; 10 minutes/solution) to 100% ethanol (two changes, 10 minutes each). The specimens were then cleared in xylene (two changes, 5 minutes each), infiltrated with two changes of paraffin (63°C, 1 hour each) and embedded in paraffin blocks. Sections were cut at 8 μm and collected after floating in a warm water bath onto prepped slides (Superfrost Plus, Fisher Scientific). Sections were deparaffinized and rehydrated (two changes of xylene, 100% and 95% ethanol,
5 minutes each, followed by 5 minutes in 75% ethanol, two changes of water, and mounted in 1:1 glycerol/PBS. Slides were observed and photographed using a Nikon Eclipse E800 photomicroscope and an Olympus IX81 confocal microscope.

In situ confocal microscopy.

Whole mount in situ hybridization was conducted using methods described previously (Tucker, 2010; based on methods found in Martindale et al., 2004). Antisense and sense NtVSP 168100 digoxigenin-labeled riboprobes were made using reagents supplied by Boehringer Mannheim; this was also the commercial source of the anti-digoxigenin and the colorization substrates.

For qPCR three animals per sample were pooled in a 1 ml homogenizer, lysed with 500 μl of Triton and homogenized. The lysate was sheared through a 26 gauge needle until little to no resistance was felt though the syringe. Samples were then normalized on ice for 15 minutes followed by 15 minutes at room temperature. 100 μl of chloroform was added to each sample, which was then shaken vigorously for 15 seconds and allowed to separate at room temperature for 5 minutes. The sample was then allowed to precipitate at room temperature for 10 minutes, then centrifuged at 14 k for 15 minutes at 4°C. The RNA pellet was rinsed with 1 ml of 70% ethanol and then spun at 14 k for 5 minutes. The resulting pellet was air-dried and suspended in RNase-free water. The quality and quantity of RNA was determined by gel electrophoresis and a Nanodrop spectrophotometer. One μg of total RNA was reverse transcribed into cDNA by incubating for 1 hour at 37°C using AMV Reverse Transcriptase (Affymetrix) and oligo dT primers. Three cDNA replicates of each treatment (control or regenerating) were used to conduct the qPCR. The composition of the PCR mix for each sample was as follows: 10 μl of cDNA template, reverse primer (1 μM 5'-GTCCTGAAACTGACCG-AGTGTCC-3'), forward primer (1 μM 5'-CAAGAAGGCTGGGACAGATGCAGTGC-3') and 12.5 μl of SybrGreen I Master Mix (Applied Biosystems). Control primers were based on 18S rRNA sequences (Reitzel and Tarrant, 2009) (5'-GACTCAACACGGGGAAACTC-3' and 5'-GCACACCACCATAGATCC-3'). The amplification program was as follows: an initial denaturing step at 95°C for 15 seconds and allowed to separate at room temperature for 5 minutes.


Competing Interests

The authors have no competing interests to declare.

References


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