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Characterization of Hydra Type IV Collagen

TYPE IV COLLAGEN IS ESSENTIAL FOR HEAD REGENERATION AND ITS EXPRESSION IS UP-REGULATED UPON EXPOSURE TO GLUCOSE*

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†The abbreviations used are: ECM, extracellular matrix; BM, basement membrane; HM, hydra medium; LEP, localized electroporation; UTR, untranslated region; bp, base pair(s); kb, kilobase(s); CHAPS, 3-[3-cholamidopropyl]dimethylammonio]-1-propanesulfonic acid; FITC, fluorescein isothiocyanate; contig, group of overlapping clones.

**The nucleotide sequence(s) reported in this paper has been submitted to the GenBank®/EMBL Data Bank with accession number(s) AF282902.

The mesogelea is a 0.3–3-,μm-thick sheet of extracellular matrix that extends throughout the freshwater coelenterate *Hydra vulgaris*, with the exception of the mouth and the aboral pore (1). In 1961, Fawcett (2) first noticed the physical similarities between mesogelea and vertebrate basement membrane, the specialized sheets of ECM composed mainly of type IV collagen, laminin, nidogen/entactin, and heparan sulfate proteoglycan that underlie or surround cells. Although the mesogelea lies between two organized epithelial cell layers and has an obvious role in physical support of the organism, this ECM is also implicated in several cellular processes. In *vitro* studies showed that epithelial cells (3) and nematocytes (4) could attach to and migrate along isolated mesogelea or substrata coated with purified ECM components such as collagen IV and laminin. Zhang and Sarras (5) proposed that in addition to cell-cell interactions and chemotactic gradients, cell migration also depends on cell-matrix interactions. Using two modified grafting procedures, in which normal cell-matrix interactions were perturbed, they showed that interstitial cell migration is sensitive to alterations in collagen structure and that migration is inhibited by molecules that can compete with cell-fibronectin interactions.

The relation of mesogelea to higher metazoan connective tissue has been speculated on for at least the past century. By the early 1950s it was recognized that mesogelea was structurally and chemically similar to vertebrate connective tissues. Since then several biochemical studies have provided evidence of collagen-like molecules in isolated hydra mesogelea (6–8). More recent studies on mesogelea by Sarras et al. (9, 10) using indirect immunocytochemistry suggested the presence of molecules similar to type IV collagen, laminin, and heparan sulfate proteoglycan core protein (all characteristic of vertebrate BM) in mesogelea, as well as fibronectin, a component of interstitial ECM. In frozen sections, antibodies to ECM components appeared localized to the mesogelea layer between the epithelial bilayer. The presence of a hydra laminin in mesogelea was confirmed when cDNA clones encoding a β1 chain of laminin were isolated (11). A fibrillar collagen has also recently been isolated from mesogelea and characterized (12), providing further evidence that mesogelea represents a primitive ECM exhibiting composite basement membrane and interstitial matrix properties.

So far six genetically different mammalian type IV α chains, α1(IV) to α6(IV), have been identified which interact to form various heterotrimeric triple-helical isoforms (13, 14). Type IV α chains have also been characterized in invertebrates including *Drosophila, Caenorhabditis elegans*, sea urchin, and sponge (15–19). The primary structure of all the α(IV) chains is similar, each chain having a short noncollagenous sequence at the N terminus, a central collagelous region consisting of Gly-X-Y repeats, and a highly conserved C-terminal noncollagelous
Isolation and Characterization of cDNA Clones—ECM-enriched segment hydram were prepared (9) and allowed to regenerate for 24 h. Poly (A+) RNA was extracted and used as template for cDNA synthesis with either oligo(dT) or random primers. The random primed cDNA was cloned into Stratagene’s ZAP II vector (the HZAPII library), whereas the 3’-biased oligo(dT)-primed cDNA was directionally cloned into UniZAP vector (the UHAPII library). Recombinant clones of the HZAPII library were screened with a 3’-poly(A)-labeled 2.3-kb hydra fibrillar collagen probe, HCl, which encodes the 5’ half of hydra type I/II collagen (12) and washed first at low stringency (45 °C, 0.2× SSC, 0.1% SDS) then at high stringency (65 °C, 0.1× SSC, 1× SDS). After each wash, filters were exposed to x-ray film for 16 h. Plaques that hybridized only at low stringency were chosen for secondary and tertiary screening. In situ excised (ZAPPEd) clones were cycle sequenced using an ABI PRISM XL 377 DNA sequencer in combination with a Big Dye terminator chemical kit (PerkinElmer Life Sciences). The DNA clones were sequenced in both directions using a primer walking strategy shown in Fig. 1. Comparisons of DNA sequences were conducted using the Blastn and Blastx programs and the GenBankNCBI database (NCBI, National Institutes of Health). Protein alignments were obtained using the Blastp, GCG, and MacVector 5.0

Isolation and Analyses of Type IV Collagen from Hydra Mesoglea—Hydra Culture—H. vulgaris were maintained at 18 °C in hydra medium (HM) as described previously by Sarras et al. (9). Cultures were washed regularly and fed Artemia salina (brine shrimp) three times a week.

To remove cell surface proteins still bound to the mesoglea, the EDTA extract was made 1% in mercaptoethanol and separated by reversed phase chromatography on a Vydac C-18 column (Separations Group, 250 × 2.2 mm) at a flow rate of 0.2 ml/min using a gradient of 5–70% acetonitril in 50 min. Two major peaks at 21 and 35% of the organic modifier were obtained, the latter of which contained the type IV collagen. After lyophilization, the isolated chains were dissolved in 70% formic acid or 0.2 M mercaptoethanol, centrifuged at 4000 rpm in a Beckman 50 Ti rotor. The supernatant was dialyzed against 20 mM Tris buffer, pH 7.4, and separated on a Hitrap Q column (1 ml) using a gradient of 0–1 M NaCl in 40 mM. Hydra collagen IV eluted as a broad peak in the range of 0.2–0.4 M NaCl. Samples were analyzed on 4–10% SDS-polyacrylamide gels. For electron microscopy, samples (about 50 µg/ml) were mixed with an equal volume of glycerol and, after spraying onto mica discs, rotary shadowed with carbon/platinum (23). The replicas were viewed with a Philips CM 12 electron microscope.

In Situ Hybridization—Whole mount localization of mRNA was performed using a digoxigenin-labeled RNA probe generated from clone 194.1 including the most 3’ 2505 bp presented in Fig. 2. The probe is made by EcoR I linearization and T3 promoter transcription of the entire insert including 207 bp of the open reading frame and 427 bp of the 3’ untranslated sequence. Fixation, processing, and visualization of the riboprobe in whole mount preparation was performed as described previously by Gresn et al. (24, 25). Briefly, hydra were fixed with 4% paraformaldehyde after relaxation of the polyps with 2% urethane. Specimens were subsequently treated with ethanol and proteinase K to facilitate diffusion of the probes into the epithelial bilayer. To stabilize digested tissues, specimens were refixed with 4% paraformaldehyde and then prehybridized in hybridization solution (50% formamide, 5× SSC, 1× Denhardt’s, 200 mg/ml tRNA, 0.1% Tween 20, 0.1% CHAPS, 100 mg/ml heparin) to block nonspecific hybridization sites. This was followed by a 48 h hybridization with the digoxigenin-labeled RNA probe and a subsequent wash in hybridization solution and 1× SSC. Specimens were then washed in MAB (100 mM maleic acid, 150 mM NaCl, pH 7.5) and preblocked (2–6 h) in MAB with 20% sheep serum and 1% bovine serum albumin. This was followed by the 16-h incubation at 4 °C in the same solution with anti-digoxigenin antibody (1:2000). Animals were washed eight times with MAB and then briefly in alkaline phosphatase buffer (100 mM Tris-HCl, pH 9.5, 50 mM MgCl2, 100 mM NaCl, 0.1% Tween 20). Specimens were then stained with BM purple (Vector) as substrate and incubated with 100 µM nitro blue tetrazolium and 5-bromo-4-chloro-3-nitrophenyl phosphate, purple alkaline phosphatase substrate (Roche Molecular Biochemicals), dehydrated with ethanol, and mounted in eupal (Asco Laboratoris). Functional Analysis of Hydra Collagen IV Using Localized Electroporation and Thio-oligonucleotide Antisense Constructs—Because transfection approaches have not been successfully applied to Cnidarians, we developed a procedure to specifically test the effect of antisense constructs on head or foot regeneration in hydra. This approach utilizes a localized electroporation technique (LEP) in which oligonucleotides are introduced into the apical or basal pole of hydra by a localized electroporation technique (LEP). This was used to test the hypothesis that de novo biosynthesis of hydra collagen IV is required for normal head morphogenesis following decapitation. Based on the work of Dr. Richard W. Wagner (29–31), a series of 20-mer oligonucleotides with phosphorothioate linkages were designed. Six oligonucleotides were synthesized to include four antisense sequences to portions of the 5’-UTR, initiation site, coding sequence, and 3’-UTR; a sense strand of the initiation site, coding sequence, and 3’-UTR, and a mismatch construct (the sequence of these molecules is given in the legend of Fig. 9). The oligonucleotides were introduced into hydra cells using electroporation (Bio-Rad Gene Pulser) with a micropipette drawn on a pipette puller. The ends of the micropipettes brought in contact with the hydra were polished with a micro-forg. Because hydra collagen IV is expressed in the ectoderm layer of cells, LEP was performed on the surface of the organism at the apical pole. Based on preliminary experiments, we found that decapitation had to be performed 2–4 h after LEP. This was necessary because if decapitation was performed prior to electroporation, an extensive loss of electroporated cells occurred at the cut ends of the polyp. To confirm the uptake of the oligonucleotides within the area where the electroporation was performed and also to reduce movement of the animal during LEP, hydra were chilled in a 10% heptanol solution (heptanol in hydra medium) for a maximum of 1 h. A 100 µm stock solution of 20-mer thiolated oligonucleotide (Genset Corp.) was mixed with FITC-dextran with a M, 10,000 (Molecular Probes, Eugene, OR) in a ratio of 3:1

(NC1) domain (20). Unlike fibrillar collagens, the long collagenous domain is interrupted at several sites by short, noncollagenous, sequences that are thought to impart flexibility on the molecule (21). Type IV collagen molecules interact to form a complex, irregular network that provides the BM with a high level of stability.

Zhang et al. (22) produced hydra cell aggregates and exposed the regenerating hydra to levels of glucose that mimic those in human diabetic patients (15 mm). Regeneration occurred as normal although within 72 h, glucose-treated reaggregating animals synthesized mesoglea twice the normal thickness, suggesting that the hydra may be a nonmammalian and temporally rapid model for the study of the much longer term glucose-induced basement membrane thickening seen in diabetic microangiopathy.

In this study, using a hydra fibrillar collagen cDNA as a probe in a low stringency screen, overlapping clones were isolated that encode a complete hydra collagen IV α chain. In addition, type IV collagen protein was purified from hydra mesoglea and partially sequenced. All of the peptide sequences obtained are present within the primary sequence deduced from the cDNAs. The importance of collagen IV in hydra development is demonstrated using antisense thio-oligonucleotides, which block head regeneration. In addition, we show that the mesoglea of adult (rather than the previously demonstrated regenerating) hydra is rapidly thickened and collagen IV expression is up-regulated upon exposure to glucose.

EXPERIMENTAL PROCEDURES

Hydra Culture—H. vulgaris were maintained at 18 °C in hydra medium (HM) as described previously by Sarras et al. (9). Cultures were washed regularly and fed Artemia salina (brine shrimp) three times a week.
The NC1 domain is indicated by the first shaded box, the large collagenous domain is indicated by the long box interrupted by 25 short noncollagenous sequences, and the 3’ NC1 domain is indicated by the final shaded box.

(typically 6 µl of DNA + 2 µl of FITC-dextran). After fitting the micropipette over the microelectrode (World Precision Instruments, Inc.), the oligonucleotide/FITC-dextran mixture was loaded into the micropipette. The Bio-Rad gene pulser was set at 100 ohms, 25 pulses at 100 µl of solution D, and 100 µl ice-cold isopropanol was added. The tube was incubated for 30 min at 4 °C and spun for 10 min at 10,000 × g. The pellet was resuspended in 100 µl of solution D, and 100 µl ice-cold isopropanol was added. The tube was incubated for a further hour at 70 °C and centrifuged (10,000 × g, 5 min), and the pellet was washed with 70% ethanol to remove any residual salt. After air drying, the pellet was resuspended in 10–20 µl of diethylpyrocarbonate-treated water. The concentration and purity of the RNA was determined by measuring A$_{260}$ nm and A$_{280}$ nm. Northern blotting and probing were performed as described previously (32).

RESULTS

Characterization of Type IV Collagen—The previously described hydra fibrillar type II collagen cDNA clone HCl1 (12) was used to screen a regenerating hydra cDNA library at low and high stringency. A clone, G4.1.2, was identified that hybridized to the fibrillar collagen probe only under conditions of low stringency. The clone was purified and found to contain 1.7 kb of interrupted collagenous sequence. Using an EcoRI restriction fragment of G4.1.2 to rescreen the library, several overlapping clones were isolated and purified that together cover the entire coding sequence of a H. vulgaris type IV collagen chain. The relationship of these clones to each other is shown in Fig. 1. The complete nucleotide sequence of hydra type IV collagen chain together with its deduced amino acid sequence is shown in Fig. 2. There is a 5’-untranslated region of 159 bp, an open reading frame of 5,169 bp, and a 504 bp 3’-untranslated region. The deduced translation product of 1723 residues with a predicted molecular weight of approximately 160,000 comprises a 24-residue putative signal peptide, a short 5’ noncollagenous domain of 16 residues, a 1455-residue collagenous domain, and a 228-residue 3’ NC1 domain.

The primary sequence and domain sizes (Table I) are similar to other reported type IV collagen chains. The locations of three-quarters of the interruptions in the hydra collagenous domain are conserved across species (data not shown). As in other species, twice as many long imperfections (5 or more residues) occur in the N-terminal half of the hydra type IV collagen domain. The hydra type IV collagen chain also has the single N-linked glycosylation site (Fig. 2, amino acids 122–124) conserved in other species and type IV chain types. Cysteine residues are particularly highly conserved. The 4 cysteines in the 16-residue 5’ noncollagenous domain (Fig. 2, amino acids 25–40) are also found in all other type IV chains and are required for N-terminal association of collagen IV molecules into tetramers to form the ‘7S domain’. There are five cysteine residues in the triple helical domain, mostly located in imperfections, with four being in the N-terminal half of the collagenous domain. This is comparable with other vertebrate and invertebrate...
chains (Table I). The C-terminal NC1 domain of hydra is the most conserved region of the molecule (Table I). The number and relative position of the cysteine residues in the NC1 domain is identical to all other type IV chains (Fig. 3) regardless of species.

The hydra sequence is slightly more closely related to the \( \alpha_1 \) family (\( \alpha_1, \alpha_3, \) and \( \alpha_5 \)) than to the \( \alpha_2 \) (\( \alpha_2, \alpha_4, \) and \( \alpha_6 \)) family of collagen IV chains based on: (i) the levels of sequence identity across species in the NC1 domain (Table I and Fig. 3); and (ii) the presence of the highly conserved \( \alpha_1 \)-like sequence GC-NGTK (Fig. 2, residues 122–127) in the 7 S region, whereas in \( \alpha_2 \)-like chains, the Lys is replaced by Arg. Accordingly, we designate the reported sequence hydra \( \alpha_1(IV) \) collagen.

Northern analyses revealed that hydra RNA contains two \( \alpha_1(IV) \) mRNA species, the more abundant mRNA being approximately 7 kb and the lesser, 6 kb (see Fig. 11). The most 3' clone we sequenced contained a poly(A) addition signal approximately 10 bases upstream of a poly(A) tail and completes a 5.9-kb contig. It is probable that a second poly(A) addition signal lies 3' and accounts for the longer mRNA transcript as is

FIG. 2. Nucleotide sequence of full-length cDNA, as shown in Fig. 1, and deduced amino acid sequence of \( H. vulgaris \) \( \alpha_1(IV) \) polypeptide. The overlapping cDNAs correspond to a 5855-nucleotide mRNA, including 159-nucleotide 5'-UTR and 521-nucleotide 3'-UTR. Numbering begins at the start codon. The signal peptide is yellow, and the 5' noncollagenous domain is underlined. Cysteine residues are red, RGD sequences are double underlined, and the potential N-linked glycosylation site is in light blue. Imperfections in the triple helical domain are in green, and the 3' NC1 domain is blue. The termination codon (taa) is in bold type. Peptide sequences obtained from purified type IV collagen are underlined with a dotted line.

Hydra Type IV Collagen
the case in several other collagen IV genes (20, 33).

Molecular and Supramolecular Structure—Extraction of hydra mesoglea with EDTA, 1 M NaCl solubilized two major collagen with molecular masses of 155 and 290 kDa (Fig. 4). The smaller component has recently been shown to represent a fibrillar collagen, hydra collagen I/II, that forms a large network of thin fibrils (12). Here we show that the larger component is type IV collagen, because all the peptide sequences obtained from the 290-kDa chain (Fig. 2) are found within the hydra a1(IV) collagen-deduced sequence. The peptide data demonstrate that the polypeptide chain is not processed by removal of the NC1 domain, as expected, and that the hydra collagen IV frequently contains hydroxyproline in the Y position of the triplet (data not shown). In many positions where lysine should be found according to the cDNA sequence, Edman degradation failed to detect any amino acid residue. This is most probably because of the presence of glycosylated lysine (or hydroxylysine) residues that are not seen in normal protein sequencing protocols.

As shown in Fig. 4, the hydra type IV collagen was not able to enter an SDS-polyacrylamide gel in the unreduced state, indicating extensive polymerization via disulfide bonds. However, even after reduction, only bands down to the size of a dimer (290 kDa) were found. Monomers were never detected, indicating that two polypeptide chains are held together via nonreducible cross-links. Most probably intermolecular bonds between two adjacent molecules are formed rather than intramolecular bonds within the same triple helical molecule.
that should result in a mixture of dimers and monomers.

To get information about the structure of the dimer we isolated type IV collagen by ultracentrifugation and ion exchange chromatography in the presence of mercaptoethanol but in otherwise non-denaturing conditions. Electron microscopy after rotary shadowing of the protein showed dimers connected via the C-terminal globular domains (Fig. 5). Surprisingly, in marked contrast to the vertebrate type IV collagens, the triple helix exhibits many bends and kinks so that the predicted length of almost 400 nm is never observed. As seen in one molecule in Fig. 5, the collagenous domain is even able to fold back on itself. The supramolecular structure of the type IV collagen in the unreduced state was visualized by electron microscopy of the unfraccionated EDTA extract. Although this fraction contains a mixture of the hydra fibrillar collagen I/II and the type IV collagen, assignment of the type IV structures was very easy because collagen I/II forms a clearly distinguishable fibrillar network lacking globular domains (12). Some selected electron microscopy pictures are shown in Fig. 6. Irregular networks are seen typically, but in addition, smaller oligomeric structures formed by lateral association of a few molecules can also be detected.

In Situ Hybridization—In situ hybridization analysis in adult hydra revealed that collagen IV expression is observed along the entire longitudinal axis of the animal but is most intense at the base of the tentacles, at the site of battery cell

### Table I

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**FIG. 3.** Amino acid sequence comparison of the NCI domain of hydra type IV collagen (Hy) with all six human type IV collagen chains (A1-A6). Dashes represent identities with the amino acid sequence of the hydra NCI domain. Gaps indicated by arrows are introduced to achieve best alignment. Conserved cysteine residues are indicated by filled arrows.
transdifferentiation (Fig. 7, A and B). Interestingly, collagen IV expression is restricted to the ectoderm (Fig. 7, C and D), whereas laminin β1, the other basement membrane component characterized in hydra, is expressed solely by the endoderm (see “Discussion”). During head regeneration type IV collagen levels are increased basal to the decapitation site. Initially, an apical cap of type IV-positive cells form (Fig. 8, A–C), and then, by 72 h, additional patches of stronger expression appear in the tentacle region (Fig. 8D).

**DISCUSSION**

Although the presence of type IV collagen in hydra had previously been suggested by a study using antibodies raised against vertebrate BM components (9, 10), the elucidation of the primary structure of collagen IV (Figs. 1 and 2) provides the first molecular identification of this protein in hydra. Sequence
The restriction of hydra type IV expression to the ectoderm from purified protein (Fig. 2) originates from the α1(IV) chain. Solubilization of type IV collagen from vertebrate basement membranes usually requires pepsin digestion, and it was therefore very surprising that the hydra type IV collagen could be solubilized in the form of dimeric molecules under reducing but otherwise nondenaturing conditions (Figs. 4 and 5). Sequence analysis of peptides obtained by harsh protease treatment of the insoluble residue remaining after extraction revealed that all the type IV collagen had been solubilized (data not shown), thus excluding the possibility that solubilized material is only a minor fraction. Electron microscopical investigation (Fig. 6) showed that the dimers assemble into a tight network stabilized by disulfide bonds. For vertebrate type IV collagen a model has been proposed in which four molecules aggregate via their N-terminal domains forming spider-like structures. The interactions are stabilized via disulfide bonds and lysine-derived cross-links, resulting in a highly protease-resistant 7 S domain that can easily be isolated after pepsin digestion. In addition, the C-terminal globular domain, NC1, binds to itself, mainly via disulfide bridges, to form a linear dimer (14). Both interactions at the N-terminal and C-terminal ends lead to the proposal of an open network structure that can further polymerize via lateral aggregation of the triple helical domains (34). Our data certainly support the idea of lateral aggregation of type IV molecules, which is very difficult to observe experimentally in vertebrate basement membranes, although there are also marked differences in the molecular architecture. Firstly, two molecules of hydra collagen type IV are held together by nonreducible cross-links between the globular NC1 domains rather than by disulfide bonds (Fig. 5). Secondly, we do not have any structural evidence for the existence of a 7 S domain in hydra collagen IV despite the presence of the 4 conserved cysteine residues in this region. Pepsin digestion of the hydra type IV collagen did not yield the 7 S domain but instead resulted in a complete destruction of the protein; a protease-resistant domain could not be detected by SDS gel electrophoresis or by electron microscopy (data not shown). It is interesting to note that lack of a 7 S domain is not unique to hydra because similar results have previously been obtained for the collagen type IV molecule of the helminth Ascaris suum (35). Thus, formation of compact, protease-resistant 7 S domain might be an important step in the polymerization of vertebrate collagen type IV but appears not be essential for invertebrates. For hydra, a less compacted and highly flexible organization of the collagen IV network is presumably advantageous given the organisms physiological requirements for an extremely flexible mesoglea.

The hydra is a member of the phylum Cnidaria, one of the oldest metazoan phyla and the highly conserved nature of type IV collagen indicates its critical role in hydra ECM formation. The presence of both collagen IV and laminin in hydra and their co-localization in the mesoglea (11) provide compelling evidence that the mesoglea is essentially composed of the same molecular components found in vertebrate BMs, and it is likely that cell-substrate interactions involve the same ligands and receptors. However, it is noteworthy that in addition to classical BM components, the hydra mesoglea also contains a centrally located core of another collagen, which, based on sequence comparisons, is a classical fibrillar collagen (12). It is therefore possible that the first functional extracellular matrices evolved with composite properties of what are now considered classical basement membrane and interstitial matrices and that later in evolution, the uses of components in these “primitive basement membranes” were refined to produce the wide array of connective tissues seen in higher orders.

The restriction of hydra type IV expression to the ectoderm

Fig. 7. Expression pattern of type IV collagen in adult hydra. Whole mount in situ hybridization using an RNA probe generated from clone 194.1 is shown. Localization of type IV collagen is observed along the entire longitudinal axis of the polyp (A) but is most intense at the base of the tentacles (B). The localization is restricted to the ectoderm (A and B). This is best seen in thin sections of whole mounts embedded in JB4 compound as shown in C and D. A bright field image is shown in C with the ectoderm containing reaction product for type IV collagen mRNA indicated by an arrowhead. The endoderm is negative and is indicated by an asterisk (C). The intervening ECM (arrow) is shown using Normarski optics in D. The ectoderm (arrowhead) and endoderm (asterisk) are also shown in D. Bar magnifications: A, 500 μm; B, 100 μm; C and D, 20 μm.

analysis clearly shows the molecule is similar to all known α(IV) chains but most closely resembles vertebrate and invertebrate α1(IV) collagen chains (Table I and Fig. 3).

Our data suggest that the hydra type IV collagen molecule is homotrimeric because all the peptide sequence data obtained
(Fig. 7) contrasts with laminin, which is expressed solely in the endoderm but is localized to the subepithelial zones adjacent to both the ectoderm and endoderm layers (11). In an immunocytochemical study of hydra ECM, polyclonal antibodies generated against mammalian type IV collagen were localized throughout the entire mesoglea (9, 10), suggesting that, like laminin, type IV collagen can assemble into basement membranes on cells that have not produced it. The sources and final locations of type IV collagen have been determined in some vertebrate and invertebrate systems. In co-cultures of fetal intestinal chick mesenchyme with rat endoderm, type IV collagen in the resulting basement membrane was derived only from the mesenchyme (36), whereas in C. elegans type IV has been detected at sites distant from its site of synthesis (37). The assembly of type IV collagen in hydra, away from the ectoderm cells that express it, suggests there is a mechanism regulating its assembly that is directed by interaction with other (cell surface-associated) molecules. Functional antisense studies show inhibition of type IV collagen translation causes a subsequent blockage in head regeneration (Fig. 9), re-emphasizing the importance of this molecule and the mesoglea in regeneration and development.

In mammals, thickening of the basement membranes in blood capillaries is the hallmark of diabetic microangiopathy, a
severe long term complication of diabetes mellitus is the leading cause of blindness and renal failure in the developed world. The molecular mechanism(s) underlying this thickening are still undetermined, although the process is thought to be glucose-dependent. Zhang et al. (22) showed that reaggregating hydra exposed to 15 mM glucose developed a newly synthesized mesoglea twice the thickness of controls and suggested hydra to elevated glucose levels retained by cells that are freely permeable to glucose. Given the striking conservation of hydra and mammalian BM genes, together with the rapidity with which glucose-responses are elicited, the hydra represents an experimentally tractable, simple, in vivo model system in which to further investigate the mechanism underlying glucose-induced basement membrane thickening.

In summary, we have characterized hydra type IV collagen and shown it to be highly conserved with respect to its mammalian counterparts, although its organization and assembly at the supramolecular level shows some unique features. Type IV collagen expression is essential for hydra development and responsive to elevated levels of glucose.

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\(^1\) M. Sarras, Jr., unpublished observations.