Sonic hedgehog delivery from self-assembled nanofiber hydrogels reduces the fibrotic response in models of erectile dysfunction

Shawn Choe¹, Dorina Veliceasa¹, Christopher W. Bond², Daniel A. Harrington³, Samuel I. Stupp⁴, Kevin T. McVary⁵, and Carol A. Podlasek⁶,*

¹Department of Urology, University of Illinois at Chicago, Chicago, IL 60612
²Department of Allergy/Immunology, Northwestern University, Feinberg School of Medicine, Chicago, IL 60611
³Department of Biosciences, Rice University, Houston, TX 77005
⁴Simpson-Querrey Institute for BioNanotechnology, Departments of Chemistry, Department of Materials Science and Engineering, and Biomedical Engineering, Northwestern University, Feinberg School of Medicine, Chicago, IL 60611
⁵Division of Urology, Southern Illinois University School of Medicine, Springfield, IL 62794
⁶Departments of Urology, Physiology and Bioengineering, University of Illinois at Chicago, Chicago, IL 60612

Abstract

Erectile dysfunction (ED) is a serious medical condition in which current treatments are ineffective in prostatectomy and diabetic patients, due to injury to the cavernous nerve (CN), which causes irreversible remodeling of the penis (decreased smooth muscle and increased collagen), through a largely undefined mechanism. We propose that sonic hedgehog (SHH) and neural innervation, are indispensable regulators of collagen in the penis, with decreased SHH protein being an integral component of the fibrotic response to loss of innervation. We examined collagen abundance and morphology in control (Peyronie’s), prostatectomy and diabetic patients, and in rat models of penile development, CN injury, SHH inhibition and under regenerative conditions, utilizing self-assembling peptide amphiphile (PA) nanofiber hydrogels for SHH delivery. Collagen abundance increased in penis of ED patients. In rats, collagen increased with CN injury in a defined time frame independent of injury severity. An inverse relationship between SHH and collagen abundance was identified; SHH inhibition increased and SHH treatment decreased penile collagen. SHH signaling in the pelvic ganglia (PG)/CN is important to maintain CN integrity and when inhibited, down stream collagen induction occurs. Collagen increased

*Correspondence to: Carol Podlasek, Ph.D., Department of Urology, M/C 955, University of Illinois at Chicago, 820 S. Wood St., CSN 515, Chicago, IL 60612, Phone: 312-996-7955, cap325@uic.edu.

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throughout penile development and with age, which is important when considering how to treat fibrosis clinically. These studies show that SHH PA treatment reduces collagen under regenerative post-prostatectomy conditions, indicating broad application for ED prevention in prostatectomy, diabetic and aging patients and in other peripheral nerve injuries. The PA nanofiber protein vehicle may be widely applicable as an in vivo delivery tool.

Graphical Abstract

Keywords
self-assembling peptide amphiphile nanofiber hydrogels; Sonic hedgehog; penis; collagen; cavernous nerve injury

Introduction

The cavernous nerve, which provides innervation to the penis, becomes injured during prostatectomy surgery, in diabetic patients, and with aging, resulting in remodeling of penile morphology and erectile dysfunction (ED). The penis is composed of smooth muscle, collagen and elastic fibers. In response to peripheral nerve injury, smooth muscle and elastic fibers decrease while collagen increases in both ED patients [1,2] and animal models [3,4]. This is an irreversible process that underlies ED development. Aging and consumption of high fat diets are also associated with increased collagen abundance and thickening of fibers [5–7]. Current ED treatments are not effective to prevent the penile remodeling, nor long term ED development. The mechanism of how collagen induction occurs in response to CN injury is complex and remains largely undefined.

As is the case with other peripheral nerves, CN regeneration efforts have not translated into improved clinical outcomes, and there is minimal understanding of how neuronal changes impact tissue remodeling. Few factors have been identified in the penis that impact collagen induction. The most widely explored is the TGF-β pathway, which in patients, increases collagen abundance 2.5–4.5 fold [8] and in animal models increases with CN injury [9]. TGF-β-1-induced collagen synthesis is inhibited by cyclic AMP synthesis in human corpora cavernosal smooth muscle cells [10], suggesting a potential mechanism of how TGF-β may be regulated. TGF-β also functions down stream of the adenosine receptor A(2B)R which has been implicated to increase proliferation in corpora cavernosal fibroblast cells [11]. Accumulating evidence suggests that cross talk may occur between the sonic hedgehog (SHH) pathway and TGF-β signaling in several diseases including gastric carcinoma [12], melanoma bone metastasis [13] and pulmonary fibrosis [14]. It is thought that Hedgehog
may mediate epithelial-mesenchymal crosstalk in pulmonary fibrosis, with SHH inducing TGF-β in lung fibroblasts while TGF-β induces SHH in cultured alveolar epithelial cells [15]. Thus, we hypothesize that SHH may also serve as an important mediator/regulator of collagen synthesis in normal and injured penis.

We have shown in previous studies that the SHH pathway is critical for the response of the penis and of the CN to denervation, regulating both penile and CN architecture, and smooth muscle apoptosis [16–23]. The SHH pathway has recently been suggested to also play an important role in the pathogenesis of fibrosis [24,14], with activation of the pathway present in fibrotic diseases such as sclerosis, interstitial pneumonitis, injury-related inflammations [25–27] and idiopathic pulmonary fibrosis [28]. In support of this idea, during development, the expression pattern of type XVIII collagen in the ureter bud is responsive to changes in SHH expression in the epithelium [29]. In the adult, hedgehog (Hh) signaling can regulate lung fibrosis [30], and expression of SHH and GLI were correlated with cerulean-induced fibrosis in the pancreas [31], suggesting that SHH may be a regulator of collagen. In this study we will examine if collagen production is responsive to SHH signaling in the penis and in the CN, and to CN regulation, and thus may provide a novel avenue for clinical intervention post prostatectomy and in diabetic patients.

We have described the use of self-assembling peptide amphiphiles (PA) as biological delivery vehicles, to prevent ED-related smooth muscle apoptosis in the penis [20,21]. These versatile hydrogel systems can be molecularly pre-programmed for SHH protein delivery, either from (1) an injectable solution with fast, in situ assembly into a soft hydrogel, or (2) highly aligned monodomain nanofiber bundles with increased mechanical integrity [32, 33]. In both permutations, these PAs offer a customized, biodegradable, solution for delivering proteins in a controlled manner over extended periods, and are easily translatable to patients in the clinic. In this study we will examine a novel neuronal component to collagen regulation and the role of the SHH pathway in the fibrotic response to nerve injury. We will utilize these innovative PA systems for SHH delivery to the luminal surfaces of the corpora cavernosa (via in situ gelation), and SHH delivery to the injured CN from a manipulable supramolecular cable (via monodomain aligned nanofibers). These materials and their technology have potentially broad application to other peripheral nerves and the tissues that they innervate.

Materials and Methods

Animals

199 Sprague-Dawley rats postnatal day 7 (P7) through P300 were obtained from Charles River. The study was carried out in strict accordance with the recommendations in the Guide for the Care and Use of Laboratory Animals of the National Institutes of Health. The animal care protocol was approved by the Office of Animal Care and Institutional Biosafety at the University of Illinois at Chicago (OACIB) and by the IACUC committee at Northwestern University and animals were cared for in accordance with institutional approval.
Patient tissues

Corpora cavernosal tissue was obtained from 21 patients who underwent penile prosthesis implantation at Northwestern Memorial Hospital. Eight patients had a previous prostatectomy, five had diabetes, and eight control patients underwent corrective surgery for Peyronie’s disease. Peyronie’s disease is a condition of the penis characterized by the alteration in the appearance and cellularity of collagen within the tunica albuginea, which becomes fibrotic with disease progression. While Peyronie’s patients have intromission difficulty, the underlying defect is tunical so the lacunar tissue remains fundamentally normal [34]. Peyronie’s tissue is the best available control other than cadaver tissue and corporal tissue was obtained from a region away from the involved tunica. For the penile prosthesis cases, the corporal tissue at the site of corporotomy was identified, minimally handled and a small wedge of lacunar tissue excised prior to any dilation or other corporal manipulation was performed. For the control Peyronie’s cases, the corporal tissue at the site of tunical defect and the surrounding corpora in the vicinity of the planned grafting site were identified. Similarly these normal tissues were minimally handled and a small wedge of lacunar tissue excised prior to any other corporal manipulation or grafting was performed. The tissue was immediately handed to a lab technician present in the operating room and the tissue was snap frozen in liquid nitrogen or fixed in 4% paraformaldehyde overnight at 4°C prior to paraffin embedding. Exclusion criteria included patients under 18 years of age. The complete study protocol was approved by the institutional review board of Northwestern University and written informed consent was obtained from all patients. For the prostatectomy patients it had been 1–17 years since surgery with an average of 6 years. For the diabetic patients, the onset of diabetes was between 7 and 24 years with an average of 12 years.

Hydroxyproline assay

Collagen abundance was quantified on frozen corpora cavernosal tissue using a modified hydroxyproline assay [35]. Each sample was examined in duplicate. The absorbance was read at 550 nm using a plate reader and the collagen content was calculated as μg collagen/mg tissue.

Trichrome stain

Trichrome stain was performed as described previously [36] on penis tissue that was fixed in Bouin’s fixative for 48 hours at 4°C prior to staining and was quantified by Image J analysis (Image J version 1.45 s, download date 5/22/2012). After background subtraction, the total area of blue (collagen) was selected and quantified in trichrome photos independently of smooth muscle (red), which decreases with nerve injury. Total area of blue/collagen was measured in 25 photos (200X magnification) taken randomly in each tissue (5 photos per section and 5 sections per tissue).

Timecourse of postnatal collagen development

Postnatal day 7, P13, P22, P40, P62, P120 and P200–300 Sprague Dawley rat penis (n=3 for each time point) were fixed in Bouin’s fixative for 48 hours at 4°C prior to examination by trichrome stain to evaluate collagen morphology and abundance with age.
Timecourse of collagen induction after CN injury

Rats were anesthetized and the pelvic ganglia (PG)/CN were surgically exposed and microforceps (size 0.02 × 0.06 mm) were used to crush the CN bilaterally for 30 seconds (constant pressure with closed forceps), greater than 5 mm from the PG. This method of CN crush has commonly been used [37,38] and the extent and reproducibility of crush injury were previously verified [21]. Sham surgery (control) was performed by exposing but not crushing the CN (n=6). Rats were sacrificed 4, 7 and 14 days after injury (n=16). For severe crush injury, three consecutive 30 second crushes were performed in the same region of the CN (n=4) and rats were sacrificed at 4 days after injury. For resection injury, a 5 mm portion of the CN was removed 5 mm from the PG (n=4) and rats were sacrificed at 4 days after injury.

SHH inhibited penis

Affi-Gel beads (100–200 mesh, Bio-Rad, Hercules, CA) were equilibrated with 100 μl of 5E1 SHH inhibitor (378 μg/ml, Jessel, Hybridoma Bank University of Iowa) or mouse IgG in 1XPBS (control) overnight at 4°C. Approximately 100 μl beads were injected into the corpora cavernosa of the penis of each rat. SHH inhibited rats were sacrificed after 1–2 (n=6), 4 (n=3) and 7 days (n=3) in comparison to mouse IgG/PBS controls (n=6, 6 and 3, respectively).

CN injury with SHH treatment of the penis by peptide amphiphile (PA)

We have previously employed self-assembling peptide amphiphiles as hydrogel carriers for controlled delivery of SHH to relevant tissues in the penis. We have used these carriers in two variants: either through injection and subsequent gelation of V3A3E3-COOH PAs directly within the corpora cavernosa (as a soft gel, coating the interior corpora cavernosal surfaces), or external preparation of V2A2E2-NH2 PAs to form highly aligned, monodomain hydrogels with sufficient mechanical integrity that they can be manipulated and placed on top of an exposed CN. In both cases, SHH is entrapped within the PA hydrogels during cation-based assembly and crosslinking. Both systems were employed in the current work, to assess SHH impact on collagen abundance in penile tissues.

Bilateral CN cut with either SHH (n=12) or BSA (n=12) treatment was performed by injection of a self-assembling peptide amphiphile (PA) hydrogel (V3A3E3-COOH) into the corpora cavernosa of the penis as described previously [20]. A 5-mm section of the CN was removed bilaterally, 5 mm from the PG. A 20-mM solution of PA (50 μl) was added to 5 μl of a solution of SHH protein in water (1.25 μg/μl, R&D Systems, Minneapolis, MN, USA). A 40 mM CaCl2 solution (50 μl) was added to the SHH-PA solution. The skin covering the penis was then retracted and the PA was immediately injected directly into the corpora cavernosa with a 26-gauge needle (105 μl volume) where the PA formed a loose gel lining the sinuses within 30 seconds to 1 minute. The final concentration of PA was 10 mM and CaCl2 was 20 mM. The final amount of SHH protein injected was 6.25 μg per rat. Penises were harvested from euthanized males by sharp dissection 2, 4 and 7 days after SHH protein/PA/CaCl2 injection and were frozen in liquid nitrogen.
SHH protein was also delivered to the penis of CN injured rats by injection of Affi-Gel beads (100–200 mesh, Bio-Rad Laboratories, Hercules, CA) equilibrated with either 1X SHH peptide (n=10, 7.5μg per animal, R & D Systems), 2XSHH peptide (n=9, 15 μg per animal, R&D Systems), or PBS (control, n=9) overnight at 4°C. Approximately 30–40 beads were injected directly into the corpora cavernosa. CN injury only rats served as a positive control (n=4). Collagen was quantified by hydroxyproline assay 4 days after CN injury.

**SHH inhibition in the PG**

Affi-Gel beads were equilibrated with 5E1 SHH inhibitor (378 μg/ml, n=12) or mouse IgG in PBS (control, n=8) overnight at 4°C. Approximately 10–20 beads were injected directly under the PG bilaterally in adult Sprague-Dawley rats. Injection was not made into the ganglia itself since this would likely destroy the ganglia. Penis tissue was dissected after 2 days of SHH inhibition in the PG.

A second SHH inhibitor, cyclopamine (10μM, n=4) was administered to Sprague Dawley rats by Affi-Gel beads under the PG for 7 days in comparison to PBS treated controls (n=4) and penis tissue was assayed for collagen by hydroxyproline.

**SHH treatment of uninjured PG/CN**

Affi-Gel beads (100–200 mesh) were equilibrated with SHH or BSA (control) protein (0.25μg/μl) overnight at 4°C. Approximately 10–20 beads were injected under the PG bilaterally and rats were sacrificed after 2 days. Penis from SHH (n=4) and BSA treated (n=3) rats were frozen for hydroxyproline analysis.

**CN crush with either SHH or BSA/MSA protein treatment of the CN by linear monodomain PA**

(C16)-V₂A₂E₂-(NH₂)₂ PA was prepared as previously described [21]. Briefly, a 100mM solution of (C16)-V₂A₂E₂-(NH₂)₂ in water was heated to 80°C, held for 30 minutes, and slowly cooled to room temperature. 200mM CaCl₂ was added to a glass slide and 8μl of the heat-cycled PA plus either 2.27μg SHH or bovine serum albumin/mouse serum albumin (BSA/MSA, control) proteins were extruded with a pipet tip onto the slide to form a linear monodomain PA hydrogel. PG/CN were crushed bilaterally for 30 seconds as described above. Linear PA with the protein was transferred with forceps on top of the crushed CNs bilaterally so that each rat received 4.54μg SHH or BSA/MSA protein. The release rate of SHH protein from the PA was previously determined to be 90% by 75 hours [21]. Rats were sacrificed after 1–2 and 4 days. Penis from SHH (n=14) and BSA/MSA treated (n=10) rats were frozen for hydroxyproline assay.

**Immunohistochemical analysis (IHC)**

IHC was performed on control (n=3) and CN injured (n=3) frozen penis sections which were cut 14μm and were fixed in acetone at 4°C for 15 minutes. OCT was removed with two washes of 1X PBS prior to blocking with 3% milk in PBS for one hour at 4°C. Sections were incubated overnight at 4°C with rabbit polyclonal antibodies against collagen type I and type IV (Ab34710 and Ab6586 respectively, 1μg/μl, Abcam) and goat polyclonal antibodies against collagen type II alpha 1 (C-19, SC-7763, and N-19, SC-7764) and
collagen type III alpha 1 C-15 (SC-8781, Santa Cruz). Secondary antibodies were 1/150 diluted chicken anti-goat and chicken anti-rabbit (Molecular Probes). Sections were mounted using DPX Mounting media (Electron Microscopy Sciences, Hatfield, PA) and microscopy performed using a Leica DM2500 Microscope.

**Statistical Analysis**

Statistics were performed where appropriate using the Excel program (Microsoft) and the results were reported ± the standard error of the mean. A t-test was performed to determine significant differences (p ≤0.05) between two groups. For comparisons across three groups, statistics were performed by ANOVA with a Scheffe’s posthoc or Dunnett’s test using the SSPS statistical program.

**Results**

**Collagen quantification in corpora cavernosal tissue of Peyronie’s, prostatectomy and diabetic patients**

Trichrome stain was performed on corpora cavernosal tissue of Peyronie’s (control), diabetic and prostatectomy patients (Figure 1A) who had ED. Smooth muscle (red) and collagen (blue) were abundant in control Peyronie’s tissue. In comparison to controls, collagen was significantly increased 19% in diabetic (p=0.015) and 18% in the corpora cavernosa of prostatectomy patients (p=0.009, Figure 1B).

**Time course of collagen development in the penis**

In the rat, differentiation and growth of penile tissues occurs primarily in the postnatal period after birth [16]. We examined collagen morphology and abundance in the rat penis with increasing postnatal age (Figure 2). At P7, collagen was identifiable within the corpora cavernosa (Figure 2A). At P13, collagen fibers were more abundant and thicker in appearance (Figure 2A). By P22, a network of collagen fibers was more clearly defined (Figure 2A). Cavernae and trabeculae resemble the adult configuration at P40 and are exclusively of the adult type by P60 [39]. Between P60 and P120, substantial collagen growth occurs. Collagen appears even more abundant in the aged penis (P200–300, Figure 2A). Collagen abundance was assayed in one set of corpora cavernosal tissue by hydroxyproline assay. Collagen increased with age in the penis (Figure 2B).

**Quantification of collagen with CN injury**

Total collagen abundance was quantified by hydroxyproline assay in corpora cavernosal tissue of rats that underwent bilateral CN crush for 30 seconds. Collagen was not increased at four days after injury (p=0.992, Figure 3A) however collagen significantly increased 12% 7 days after injury (p=0.038), and 16% at 14 days after injury (p=0.010, Figure 3A). No change in collagen was observed at 4 days after injury when a more severe CN crush injury was performed (3 times 30 seconds) or CN resection (p=0.560, Figure 3B).
Localization of collagen subtypes in sham and CN injured rat penes

IHC analysis was performed on frozen penis sections assayed for collagen I–IV in sham (Figure 4) and CN injured 4day (Figure 5) rat penis. Collagen IV had the most pronounced staining throughout the corpora cavernosa and in the perineurium of the nerves in the nerve bundle. Collagen IV was also abundantly expressed in concentric rings surrounding the urethra, but was not identified within the urethral epithelium (Figure 4). Collagen III was mildly expressed in the corpora cavernosa surrounding the sinusoidal spaces. It was also highly abundant in the perineurium of nerves in the nerve bundle and in the basement membrane of the urethra (Figure 4). Collagen II was faintly expressed in the corpora cavernosa in the area surrounding the sinusoidal spaces but was not identified in the nerve bundle or in the urethra (Figure 4). This was verified using two different collagen II antibodies. Collagen I was moderately expressed in the corpora cavernosa in the region surrounding the sinusoidal spaces and also in between the sinuses. Collagen I was only faintly present in the nerves of the nerve bundle and moderately in the epithelium of the urethra (Figure 4). No change in collagen localization was observed 4 days after CN injury (Figure 5). Collagen II was identifiable in the urethra after CN injury, but not present in normal tissue. The staining intensity appeared more intense for collagen III and IV after CN injury (Figure 5). No change in localization of collagen I–IV were observed 14 days after injury (Data not shown).

SHH inhibition in the penis increases collagen abundance in the corpora cavernosa

Trichrome stain and hydroxyproline assay were used to quantify collagen in penis tissue that was treated with 5E1 SHH inhibitor via Affi-Gel beads injected into the corpora cavernosa for 1–2, 4 and 7 days in comparison to control rats that were treated with mouse IgG in 1XPBS. Trichrome stain showed abundant collagen (blue) and smooth muscle (red) in penis tissue of the control group (Figure 6A). There was a progressive loss of smooth muscle, as documented previously [19] with SHH inhibition, and collagen appeared increased by visual observation (Figure 6A). Quantification of collagen in the corpora cavernosa by hydroxyproline assay showed that collagen abundance increased in response to SHH inhibition by 17% (p=0.045) at 1–2 days of inhibition and by 21% at 4 days of inhibition (p=0.030, Figure 6B), indicating that collagen induction is responsive to changes in SHH signaling.

CN injury with SHH treatment of the penis by peptide amphiphile (PA) or Affi-Gel beads

The response of the penis to SHH signaling under regenerative conditions was examined in CN resected rats. A CN resection model was used to assay response of the penis to SHH treatment since it is the most severe form of CN injury. If a response is seen in this model than treatment is likely more effective in a CN crush model. Trichrome stain was performed on penis tissue from rats that underwent bilateral CN resection and were treated with either SHH or BSA (control) protein by PA injected into the corpora cavernosa of the penis. Smooth muscle and collagen were abundant in the SHH PA treated rats (Figure 7A). Smooth muscle appeared less abundant and collagen more abundant after injury in the BSA treated group. When quantified by Image J analysis, collagen was 19% lower in the SHH treated group than in the BSA treated group (p=0.002) at 2 days after injury and 7% lower
(p=0.039) at 4 days after injury (Figure 7B). By 7 days after injury, no difference in collagen abundance was apparent with SHH treatment (p=0.485, Figure 7B). SHH protein would be largely expended from the PA by 7 days as previously shown [20], while collagen levels increased 12% 7 days after CN injury.

These results were replicated using Affi-gel bead delivery of SHH protein to the penis at the time of CN resection, indicating that corpora cavernosal morphology (smooth muscle and collagen) appeared normal four days after CN injury in the presence of SHH protein (Figure 7C).

**Collagen abundance in the penis after SHH inhibition in the PG**

Since SHH signaling in the PG/CN impacts penile smooth muscle, we examined if it also impacts collagen. Total collagen was assayed in the corpora cavernosa by hydroxyproline quantification after SHH inhibition in the PG using the 5E1 SHH inhibitor. Collagen increased 17% in the penis when SHH was inhibited in the PG for 2 days (p=0.011, Figure 8A). These results were verified using a second SHH inhibitor, cyclopamine, administered for 7 days to the PG, which showed a 12% increase in collagen (Cyclopamine:3.99±0.251, Control:3.51±0.094, p=0.05).

**Collagen abundance in the penis after SHH treatment of uninjured PG**

Collagen was quantified by hydroxyproline assay in the corpora cavernosa of rats treated with either SHH or BSA (control) protein by Affi-Gel beads under the PG. Collagen abundance in the penis was unchanged with SHH treatment of normal/uninjured PG for 2 days (p=0.132, Figure 8B).

**Quantification of collagen in the penis after CN injury and either SHH or BSA/MSA treatment of the PG/CN by linear monodomain PA**

The response of the penis to SHH signaling under regenerative conditions was examined by quantifying collagen in the corpora cavernosa after CN crush and SHH or BSA/MSA (control) treatment of the CN via linear monodomain PA for 1–2 and 4 days. Hydroxyproline assay showed that collagen abundance was unaltered after 1–2 days of SHH treatment (p=0.240, Figure 8C). However, at 4 days, collagen in the corpora cavernosa was 9% lower (p=0.025, Figure 8D).

**Discussion**

Collagen is a major component of the extracellular matrix and is important for development of tissues and adult homeostasis [40]. There are 21 different types of collagen that have been identified in mammals, which provide strong supportive extracellular scaffolds. As the fibrillar procollagen is secreted from cells, proteases cleave the propeptides from the collagenous domains [40]. The fibrillar forms of collagen are types I, II, III, V and XI. Type I collagen is most abundant in the body, present in bones and teeth, while type II collagen composes the major structural protein of cartilage [40]. In rats, collagen is typically transversely oriented in bundles and is the most abundant component of trabeculae with a volumetric density of 62.7% [41]. For comparison, smooth muscle has a volumetric density...
of 9.1% and elastic fibers of 4.9% [41]. Human corpora cavernosa differs from that of rats by containing smaller amounts of smooth muscle, more collagen, and the presence of fibrillar collagen and smooth muscle cell subendothelial layers [41].

In this study we examined whether collagen synthesis in the corpora cavernosa of the penis is responsive to perturbation of SHH signaling. When SHH was inhibited in normal penis, collagen increased by ~20% at 1–2 days and 4 days of inhibition. This result identifies an inverse relationship between SHH levels and collagen abundance. Collagen increases in the penis with CN injury (Figure 3); since we’ve previously shown that SHH protein decreases in the penis with CN injury [18], this suggests that SHH inhibition may play a role in collagen induction. When SHH was administered to the corpora cavernosa by peptide amphiphile nanofiber hydrogel at the time of CN injury, collagen was lower in the SHH treated rats at 2 and 4 days after injury. We’ve previously quantified the in vitro release rate of SHH protein from the PA [20]. Approximately 50% of the protein is released within 24 hours and an additional ~20% is released over the following 5 days. Since the serum half-life for SHH protein is about an hour, and 50% of the protein was released in the first day, it is not surprising that collagen induction is suppressed more in the first two days when SHH protein abundance is higher, and suppression dwindles as protein levels decline (Figure 7B). TGFβ1 induction by hypoxia, which accompanies CN injury, occurs in in vitro cultured corpora cavernosal smooth muscle cells, with a peak time of 48 hours [42], suggesting the importance of this early induction window to collagen production. These findings indicate that SHH signaling, either inhibition or treatment has a selective impact on collagen abundance in the penis, when administered directly to the corpora cavernosa. These findings suggest that for maximal suppression of collagen induction after nerve injury, such as would be needed for clinical translation to prostatectomy patients, several sequential SHH PA injections might be needed to maintain SHH levels and prevent collagen synthesis.

We examined whether SHH signaling in the PG/CN also impacts collagen abundance in the penis. These studies were performed because SHH is an integral component of normal PG/CN homeostasis and when inhibited, demyelination and axonal degeneration occur, resulting in down stream smooth muscle changes in the penis and initiation of penile remodeling [21]. SHH inhibition in the PG/CN, using two different antibody (5e1) and chemical (cyclopamine) inhibitors, increased collagen in the penis (Figure 8). This finding supports the idea that CN integrity is important to regulation of collagen abundance in the penis and indicates that SHH signaling is integral to maintain CN integrity. Since SHH protein is decreased in the PG/CN with CN injury, with a maximal decrease in the active form apparent at 7 days after injury [22], and collagen increases are first measurable in the penis 7 days after CN injury, this suggests that SHH signaling in the PG/CN may play an important role in regulating collagen abundance in the penis. When the CN was crushed, SHH treatment of the CN had no effect on collagen abundance at 2 days after injury however collagen was decreased in the presence of SHH treatment at 4 days after injury. At 2 days after injury, endogenous SHH protein is reduced ~25%, and it is plausible that added SHH protein by PA may compensate for the drop in endogenous levels, resulting in no change in neural integrity or collagen abundance. At 4 days after injury, endogenous SHH protein levels in the PG/CN are reduced ~50% [22], so addition of SHH by PA would have greater impact on nerve integrity and thus down stream collagen abundance in the penis,
even if SHH protein delivered by PA is less at 4 days than 2 days, as previously documented [21]. These results indicate that collagen abundance in the penis is responsive to SHH signaling in the PG/CN, although changes in collagen take longer and are less robust than when SHH is applied directly to the corpora cavernosa. This is in opposition to the response of penile smooth muscle, which shows more pronounced apoptosis suppression when SHH treatment occurs in the CN.

Neural innervation is required to maintain collagen homeostasis in the penis, since CN crush injury causes a progressive increase in collagen abundance with time after injury (Figure 3). This is supported by our findings that SHH inhibition in the PG/CN results in increased collagen abundance in the penis (Figure 8) since SHH inhibition damages the CN. This is a novel finding since potential neural regulation of collagen has not previously been explored. We tested whether the severity of CN injury affects the time line of collagen induction by performing a more severe crush and resection injury (Figure 3B). Neither of the more severe injuries was able to increase collagen earlier than the mild crush. Thus the integrated response of the PG/CN and penis to CN injury, results in a predictable early response of the CN, including demyelination and axonal degeneration of CN fibers. Loss of innervation leads to an early wave of primarily smooth muscle apoptosis in the corpora cavernosa that peaks at 4 days after injury [22], followed by a slower induction of collagen at 7 days after injury (Figure 3). The response of collagen to loss of neural innervation might also explain why collagen appeared more abundant at P200–P300 than P120 (Figure 2), since spontaneous CN injury occurs in ~23% of Sprague Dawley rats as they age [43].

We examined the time course of collagen and smooth muscle development in the penis. In the rat, differentiation of the penile tissues occurs primarily in the postnatal period after birth. At E19, the corpora cavernosa consists of an undifferentiated mesenchyme, cuboidal epithelium and a thin, poorly defined basement membrane that the attenuated endothelium rests on [44]. Differentiation into erectile tissue containing both lacunae and trabeculae occurs during the first week after birth [45]. At P7, collagen is identifiable within the corpora cavernosa as thin fibers. At P13, collagen fibers are more abundant and thicker in appearance with a more clearly defined network. By P14, cavernous spaces appear large, irregularly shaped and are lined by an attenuated endothelium [39]. By P22, smooth muscle has the characteristic appearance and collagen fibers appear as an interconnected mesh within the corpora cavernosa. Cavernae resemble the adult configuration by P40 and are exclusively of the adult type by P60 [39]. Between P60 and the adult (P120) penis, growth of smooth muscle and collagen occur. Once the architecture of the penis is defined in the adult, it has been thought that the morphology does not change. However collagen appears even more abundant in the aged penis (P200–300), with smaller sinusoidal lumens. Collagen growth increases throughout postnatal penile development and may continue to grow/proliferate as a natural part of the aging process as CN degeneration occurs with age. This is important when considering how we think about and treat fibrosis clinically.

In an effort to understand how collagen changes in response to nerve injury, we examined the localization of collagen sub-types I, II, III and IV in normal and CN injured rat penis. Collagen IV appeared most abundant, with staining present in the corpora cavernosa, nerve bundle (including the perineurium) and in concentric layers surrounding the urethra.
Collagen III, I and II were also present in the corpora cavernosa in descending order of observed intensity. Collagen III had pronounced staining in the perineurium, while collagen I was faint. Collagen III was localized in the basement membrane of the urethra, while collagen I was present in the columnar epithelium. Collagen II was not detectable in the urethral tissue. It is interesting that the different collagen sub-types were identified in adjacent layers of the urethra and may suggest cooperative function. The perineurium and urethral staining appeared less intense after CN injury for collagen I. Collagen II was identifiable in the urethra after CN injury but not in normal tissue, and its intensity was also increased in the corpora cavernosa with injury. The localization of collagen III and IV did not change with CN injury, however the intensity of the staining appeared higher. This is consistent with a previous report of increased collagen III after CN injury [9].

In order to understand the clinical significance of changes observed in our animal models, we quantified collagen in corpora cavernosal tissue from Peyronie’s (control), prostatectomy and diabetic patients. Collagen increased ~20% in both prostatectomy and diabetic patients relative to Peyronie’s controls. This is an important observation since collagen synthesis and remodeling appears to be an ongoing process that occurs with age and is also responsive to changes in neuronal input/signaling that may occur with injury and disease. We have shown previously in these patient tissues that SHH protein is decreased in the corpora cavernosa [23], consistent with our observations that SHH inhibition causes increased collagen abundance in the penis. While it is clear that perturbation of SHH signaling either directly in the penis or indirectly in the PG/CN, impact collagen production, further study is required to understand the mechanism of how this occurs. In previous animal studies we’ve shown that SHH may have translational potential to prevent smooth muscle changes that occur with prostatectomy [20]. The results presented here suggest that SHH treatment using monodomain gels containing PA nanofibers may have the added benefit of reducing collagen under regenerative post-prostatectomy conditions. This multi-pronged approach, which impacts smooth muscle degeneration, collagen synthesis and CN regeneration, has wide implications for ED prevention in prostatectomy, diabetic and aging patients and in other peripheral nerve injuries.

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References


Figure 1.
(A) Trichrome stain of corpora cavernosal tissue from control Peyronie’s, prostatectomy and diabetic patients (100X magnification). (B) Quantification of collagen was performed from trichrome stained tissues using Image J with background subtraction. Collagen levels were significantly higher (19%, p=0.015) in diabetic and in prostatectomy (18%, p=0.009) patients in comparison to control Peyronie’s patients.
Figure 2.
Trichrome stain of penis tissue was performed at several key time points during postnatal development (P7–P300) to show collagen morphology and development (A, 50–200X magnification). Collagen was quantified in one set of corpora cavernosal tissue from each age of penis (B). Collagen abundance increased with age.
Figure 3.
(A) Quantification of collagen by hydroxyproline assay of corpora cavernosal tissue from rats that underwent sham surgery or CN crush showed significantly increased collagen at 7 (p=0.038) and 14 (p=0.010) days after injury. (B) Quantification of collagen by hydroxyproline assay of corpora cavernosal tissue from rats that underwent sham surgery, mild or severe crush injury, or CN resection showed no change in collagen at 4 days after injury (p=0.560).
Figure 4.
Immunohistochemical analysis showing the localization of collagen I, II, III and IV in the nerve bundle, urethra and corpora cavernosa of normal Sprague Dawley rat penis. (100–200X magnification).
Figure 5.
Immunohistochemical analysis showing the localization of collagen I–IV in rat penis tissue four days after CN crush (100–200X magnification).
Figure 6.
(A) Trichrome stain of penis tissue that was treated by Affi-Gel beads with either 5E1 SHH inhibitor or mouse IgG in 1X PBS for 2, 4 and 7 days (400X magnification). Smooth muscle and collagen were abundant in control penis. There was a progressive loss of smooth muscle and collagen appeared elevated with SHH inhibition. (B) Quantification of collagen by hydroxyproline assay showed a 17% increase after 1–2 days (p=0.045) and a 21% increase (p=0.030) after 4 days of SHH inhibition.
Figure 7.
Trichrome stain of penis tissue of rats that under went CN resection and SHH or BSA treatment by peptide amphiphile delivery to the penis (250X magnification). (B) Quantification of collagen by Image J analysis showed decreased collagen in CN injured penis with SHH treatment at 2 days (19% decrease, p=0.002) and 4 days (7% decrease, p=0.039) after injury/treatment. By 7 days SHH protein had been expelled from the PA and there was no difference in collagen abundance (p=0.485). (C) SHH protein delivery by Affi-Gel bead injection to the penis showed comparable results to peptide amphiphile delivery.
with unchanged corpora cavernosal morphology 4 days after CN resection in the presence of exogenous SHH protein.
Figure 8.
(A) Quantification of collagen by hydroxyproline assay of penis tissue from Sprague Dawley rats that underwent SHH inhibition in the PG with the 5E1 inhibitor or mouse IgG in PBS control. SHH inhibition in the PG for two days resulted in a 17% increase in collagen in the penis (p=0.011). (B) Quantification of collagen by hydroxyproline assay of penis tissue from rats in which uninjured/normal PG were treated with SHH protein or BSA protein (Control) for 2 days (without crush injury), showed no change in collagen (p=0.131). Quantification of collagen in penis tissue of rats that under went CN crush and SHH or BSA/MSA treatment by linear peptide amphiphile treatment for 1–2 (C) and 4 (D) days.
showed no difference in collagen at 1–2 days (p=0.240), but a 9% lower collagen abundance at 4 days of SHH treatment (p=0.025).