Genipin-induced changes in collagen gels: Correlation of mechanical properties to fluorescence

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Abstract: Controlled crosslinking of collagen gels has important applications in cell and tissue mechanics as well as tissue engineering. Genipin is a natural plant extract that has been shown to crosslink biological tissues and to produce color and fluorescence changes upon crosslinking. We have characterized the effects of genipin concentration and incubation duration on the mechanical and fluorigenic properties of type I collagen gels. Gels were exposed to genipin (0, 1, 5, or 10 mM) for a defined duration (2, 4, 6, or 12 h). Mechanical properties were characterized using parallel plate rheometry, while fluorigenic properties were examined with a spectrofluorimetric plate reader and with a standard, inverted epifluorescent microscope. Additionally, Fourier transform infrared spectroscopy was used to characterize and track the crosslinking reaction in real-time. Genipin produced significant concentration- and incubation-dependent increases in the storage modulus, loss modulus, and fluorescence intensity. Storage modulus was strongly correlated to fluorescence exponentially. Minimal cytotoxicity was observed for exposure of L929 fibroblasts cultured within collagen gels to 1 mM genipin for 24 h, but significant cell death occurred for 5 and 10 mM genipin. We conclude that genipin can be used to stiffen collagen gels in a relatively short time frame, that low concentrations of genipin can be used to crosslink cell-populated collagen gels to affect cell behavior that is influenced by the mechanical properties of the tissue scaffold, and that the degree of crosslinking can be reliably assayed optically via simple fluorescence measurements. © 2008 Wiley Periodicals, Inc. J Biomed Mater Res 87A: 308–320, 2008

Key words: crosslinking; collagen; tissue engineering; rheology; mechanotransduction; FTIR

INTRODUCTION

It is now clear that in many tissue systems, the mechnanostructural properties of the extracellular matrix contribute to the regulation of cellular functions in addition to the mechanical functions of the tissue.1,2 To investigate these phenomena and to properly design bioartificial, tissue engineered replacements, it is frequently desired to control the mechanical properties of biomaterials. Collagen-based tissue equivalents are of special interest, largely because collagen is a primary mechnanostructural element in many connective tissues, including dermis, blood vessels, tendons, and ligaments.3-6 Additionally, collagen’s superior biocompatibility and nearly ubiquitous bioactivity have made it one of the most extensively investigated biomaterial scaffolds for engineering the aforementioned tissues, and others, including hepatic7 and neural tissues.8 It is, therefore, critical to maintain the ability to manipulate the mechanical properties of collagen gels, both to study mechanotransduction and to improve the properties of bioartificial tissues. While the properties of a collagen scaffold can be altered by merely changing the concentration of collagen monomers prior to self-assembly, thereby making a more concentrated gel, most often a crosslinking mechanism is implemented.

A variety of methods exist to crosslink collagen. In vivo, tissues are naturally crosslinked by enzymes such as lysyl oxidase9,10 and transglutaminase.11,12 However, use of these enzymes for bulk changes in mechanical properties in cultured tissue mimics is cost prohibitive. Chemical treatments with aldehydes...
are often used to preserve and stiffen tissues. However, for cell-populated collagen gels (often termed “tissue equivalents”), these treatments are highly toxic. Nonenzymatic glycation has been used to improve the mechanical properties of bioartificial blood vessels in vitro by including a reducing sugar, such as ribose, in the culture medium. However, the concentrations necessary to achieve sufficient crosslinking to significantly affect the mechanical properties in a timely manner (<1–2 weeks) are toxic, requiring longer incubations at lower concentrations. Irradiation with ultraviolet (UV) light has also been used to crosslink collagen, but has limited use in cellular tissues and tissue equivalents because of the potential for UV-mediated DNA degradation. Furthermore, UV light may crosslink thicker tissues nonuniformly. Nonenzymatic nitration, which is linked to many age-associated changes, including alterations in collagen connective tissues consistent with nitrite end products of nitric oxide, has been shown to increase type I collagen crosslinking and deplete tyrosine residues, and is not immediately cytotoxic. Nitrites can also alter the structure of other proteins and enzymes to affect their regulatory functions.

Recently, genipin, a compound extracted from the fruit of the Gardenia Jasminoides, has been shown to crosslink cellular and acellular tissues, as well as biomaterials including gelatin microspheres, alginate–chitosan composites, and poly(ethylene)-glycol hydrogels. Additionally, results suggest that genipin is cell-tolerated. For these reasons, genipin has been offered as an alternative crosslinking agent for improving the mechanical properties of bioartificial tissues.

Genipin has been found to crosslink gelatin through nucleophilic attack by primary amine groups on lysine and arginine residues on the C3 atom of genipin, subsequently embedding a tertiary nitrogen in the six-membered ring in place of an oxygen atom. We expect a similar mechanism for the reaction of collagen and genipin. In addition to crosslinking collagen and increasing mechanical strength, treatment with genipin, which is white in crystalline form and produces a clear solution when dissolved in water or saline, has two unique outcomes: (1) following crosslinking with genipin, normally opaque collagen turns blue; and (2) these crosslinks emit fluorescence at 630 nm when excited at 590 nm. Thus, genipin crosslinking generates a molecular fingerprint that may be probed optically in situ to evaluate the degree of crosslinking and, possibly, the mechanical properties of collagen. Herein, we characterize the effects of genipin exposure on the mechanical properties of acellular collagen gels, and we correlate these properties to fluorescence intensity. We examine the molecular changes during crosslinking with Fourier Transform Infrared Spectroscopy (FTIR) in situ. We also assess the cytotoxic effects of direct exposure of genipin to cells in collagen tissue equivalents. These data provide a valuable blueprint for future studies applying genipin for efficient crosslinking in vitro to evaluate mechanotransduction and to assist in the design of bioartificial tissues for a variety of tissue systems.

**METHODS**

**Collagen gels**

Type I collagen gels were prepared as previously described by mixing 20 μL 1M Hepes buffer, 140 μL 0.1N NaOH, 100 μL of 10× phosphate buffered saline (PBS), 60 μL of PBS (Invitrogen, Carlsbad, CA), and 677 μL of 3.0 mg/mL collagen (Elastin Products Company, Owensville, MO) to make a 2.0 mg/mL collagen solution. The collagen solution self-assembles into a gel upon incubation at 37°C. For mechanical testing and fluorescence studies, acellular type I collagen gels were incubated in 0, 1, 5, or 10 mM genipin (Challenge Bioproducts, Taiwan) in PBS for 2, 4, 6, or 12 h. Samples were placed on a rocker to ensure adequate diffusion and equilibration of genipin through the gel.

**Mechanical testing**

Mechanical testing was done using a Rheometrics SR-2000 parallel plate rheometer with a temperature-controlled incubation chamber set to maintain 37°C (TA Instruments, New Castle, DE). A 25-mm diameter hole was punched in a 4-mm thick layer of poly(dimethyl siloxane). Collagen solution (800 μL) was pipetted into the well and transferred to a 37°C incubator to induce self-assembly. Following gel formation, 4.8 mL of PBS with a defined concentration of genipin (0, 1, 5, or 10 mM) was added to the Petri dish and the dish placed on a rocker to ensure complete mixing. Collagen gels were incubated in genipin for a defined period of time (2, 4, 6, or 12 h), after which the solution was aspirated, and gels were rinsed generously with PBS. The gels were carefully removed with a spatula and transferred to the bottom plate of the rheometer. The top plate was lowered to a height of 0.8 mm. The dynamic storage and loss moduli of the gels were evaluated at 1% shear strain amplitude at frequencies ranging from 0.1 to 10 Hz. Three samples prepared from separate batches of collagen were tested at each combination of genipin concentration/incubation duration. The data were analyzed statistically with ANOVA with genipin concentration and incubation duration as fixed effects. Significance levels were set at p < 0.05.

**Fluorescence testing**

Changes in fluorescence intensity due to genipin crosslinking were evaluated in gels prepared in a 96-well tissue
culture plate. A 40-μL aliquot of collagen was pipetted into each well. The plate was incubated at 37°C to induce self-assembly. PBS (240 μL) with defined concentrations (0, 1, 5, and 10 mM) of genipin was added to each well and the plate placed on a rocker plate to ensure equilibration of genipin throughout the gel. The gels were incubated in genipin for defined durations (2, 4, 6, 12 h) that matched the conditions from the rheology studies. At the appropriate time point, the genipin solution was removed, and the gels were rinsed extensively with PBS.

Genipin-induced fluorescence was evaluated in two ways. Some plates were transferred to the computer-controlled stage of an Olympus IX81 inverted microscope (Olympus, Melville, NY) to evaluate the feasibility of evaluating the fluorescence with standard epifluorescence microscopy for tissue engineering and mechanotransduction applications. An image of the fluorescence intensity of a representative field from each well (generally near the volumetric centroid of the gel) was captured digitally (Hamamatsu ORCA, Hamamatsu City, Japan) (590 nm Exc, 630 nm Em), and the mean intensity of the field was measured using Olympus Microsuite software (Olympus, Melville, NY). Identical exposure settings were used for all epifluorescent imaging. Each combination of genipin concentration/incubation time was tested in at least triplicate from at three replicates per condition per experiment. Separate plates were read with a Cytofluor spectrofluorimetric plate reader (Applied Biosystems, Foster City, CA) with 590-nm excitation and 645-nm emission filters to demonstrate the ability to rapidly screen the degree of crosslinking based on fluorescence.

**FTIR characterization**

The reaction between genipin and collagen was monitored in situ using FTIR for up to 4.5 h in an attenuated total reflection (ATR) geometry (Fig. 1). A type I collagen solution was pipetted on a silicon plate (~1 cm × 1.5 cm) with the longer side beveled at a 45° angle for entry and exit of the IR beam. The plate was sandwiched between two pieces of Teflon® with the top piece hollowed out to contain the collagen. The ATR setup was maintained at 37°C in a nitrogen-purged Magna-IR 760 FTIR Spectrometer (Thermo Electron Corporation, Wal-tham, MA) to facilitate self-assembly of the collagen. After self-assembly, a solution of 10 mM genipin was deposited on top of the collagen and allowed to diffuse into the gel to the silicon–collagen interface. Only the highest concentration of genipin was studied with FTIR to see the most exaggerated response to crosslinking. The infrared beam entered the silicon wafer and was reflected internally (~8x in the top face) creating an evanescent wave that probed a depth of ~2 μm above the surface of the silicon wafer into the collagen gel. Spectra for collagen crosslinked with 10 mM genipin for 12 h (with genipin solution equilibrated throughout the gel as described earlier), and then rinsed extensively with PBS to remove all free genipin, which represents the most extreme condition characterized rheometrically and fluorimetrically, and the spectra for a pure 10 mM genipin solution were similarly acquired. Spectra from untreated type 1 collagen gels served as the reference for the crosslinked collagen, while the spectrum from water was used as the reference for the genipin solution.

**Cytotoxicity**

Cytotoxic effects of genipin were evaluated using Calcein-AM (Invitogen Corp, Eugene, OR) as an indicator of live cells. L929 fibroblasts were uniformly suspended in collagen solution at 50,000 cells/ml. Aliquots of collagen solution (40 μL) were then pipetted into individual wells of a 96-well plate, which was transferred to a 37°C incubator to allow the gel to self-assemble. Genipin (0, 1, 5, or 10 mM) was added to the media, and the plates were placed on a rocker to facilitate mixing. Gels were incubated in media with genipin for 12 or 24 h. At the appropriate time point, gels were rinsed in PBS, and 20 μL of 8 μM Calcein solution was added to each well. The plates were transferred to the computer-controlled stage of an Olympus IX81 inverted microscope operating in epifluorescence mode (480 nm Exc, 535 nm Em). Three representative areas from each well were imaged serially through the thickness of the gel. The images were stacked to project all of the cells through the imaged volume onto one plane, and the cells were counted manually. The number of cells was compared across conditions with ANOVA (p < 0.05).

![Figure 1](image.png)  
**Figure 1.** ATR setup showing genipin deposition for time-resolved study. Genipin solution (10 mM) is deposited on top of and diffuses through the collagen gel. A 2-μm region of the collagen gel is probed by the IR evanescent field during genipin-mediated crosslinking. The genipin solution (10 mM) and collagen crosslinked with genipin for 12 h are probed in a similar way.
RESULTS

Mechanical testing

Rheological testing with parallel plate rheometry revealed that incubation in genipin increased the storage and loss moduli of acellular collagen gels (Figs. 2 and 3, respectively). No shrinkage of the gels was observed with crosslinking (data not shown). Storage moduli increased gradually with frequency for all conditions, and then dropped off at higher frequencies for many samples. Inspection of gels revealed damage to the samples, which did not occur if experiments were run only at lower frequencies (data not shown), and we assumed that the damage was responsible for the apparent decrease in stiffness. In general, increased crosslinking delayed this damage. Loss modulus decreased gradually with frequency in all conditions, and generally began to increase concurrent with the decrease in storage modulus, which we again attribute to damage to the gel, though the increase in loss modulus was more gradual than the corresponding decrease in storage modulus. Increasing genipin concentration and the duration of incubation also produced significant increases in storage and loss moduli ($p < 0.001$). Cell-induced strain of tissue equivalents, such as the strains produced during cell-mediated gel compaction or cell migration, generally occurs at a low strain rate.\textsuperscript{31} We therefore focused on storage moduli at 0.1 Hz, which are shown for the different genipin concentrations and incubation durations in Figure 4. Post hoc analysis (Fisher’s least significant difference test) revealed significant differences among all pairwise comparisons for the effects of genipin concentration on storage modulus (all $p < 0.001$) and all pairwise comparisons of loss modulus (max $p = 0.046$). For incubation duration, all pairwise comparisons of storage modulus were significantly different.

![Figure 2](image-url). Storage moduli following parallel plate rheometry. A: 2-h incubation; (B) 4-h incubation; (C) 6-h incubation; (D) 12-h incubation. Samples were subjected to 1% shear strain amplitude over a range of frequencies. Results are average ± std err. Both genipin concentration and the duration of incubation significantly affected the storage modulus. Storage modulus tended to decline at higher frequencies, which was associated with damage to the gels.
Similar results were obtained for pairwise comparisons of the effects of duration on loss modulus at 0.1 Hz: all pairwise comparisons were significantly different (max $p < 0.013$), except 4 h versus 6 h ($p = 0.913$). Nearly identical results were observed for comparisons of storage moduli at 2 Hz, which represents a loading rate more consistent with functions of many load-bearing tissues. Comparisons of loss moduli at 2 Hz showed significant differences between all concentrations (max $p = 0.003$) except 0 mM versus 1 mM ($p = 0.270$). Loss moduli at 2 Hz were significantly different only between 12 h and each of the other durations (max $p < 0.001$).

### Fluorescence testing

Incubation of acellular collagen gels in genipin caused the normally opaque, nonfluorescing gels to turn blue and emit a red fluorescence. The fluorescence intensity of collagen gels was measured in separate samples in parallel to the mechanical testing (Fig. 5). Fluorescence intensity measured from digital images captured with epifluorescence microscopy increased significantly with genipin concentration ($p < 0.001$) and incubation duration ($p < 0.001$) (two-way ANOVA). Post hoc analysis (Fisher’s LSD) revealed significant differences among all pairwise combinations of concentration (all $p < 0.001$) and durations (max $p = 0.003$). Similar statistically significant trends were observed in measurements taken spectrofluorimetrically ($p < 0.001$). Post hoc analysis of plate reader fluorescence revealed significant differences (max $p = 0.033$) among all pairwise comparisons of concentrations except 5 mM versus 10 mM ($p = 0.199$) and among all pairwise comparisons of duration except 4 h versus 6 h ($p = 0.240$). As with any fluorimetric (or colorimetric) optical assay, intensity measurements tended to saturate at high levels of fluorescence for both systems of measurement using a constant exposure setting.

![Figure 3. Loss moduli following parallel plate rheometry. A: 2-h incubation; B: 4-h incubation; C: 6-h incubation; D: 12-h incubation. Results are average ± std err. Loss modulus tended to decline with frequency and then rise concurrent with the decline in storage modulus at higher frequencies. Both genipin concentration and the duration of incubation significantly affected the storage modulus.](image)
The concurrent increase in fluorescence intensity with crosslinking presents a unique opportunity to assay the stiffness of the gels optically, if the fluorescence measurement can be appropriately calibrated against a measure of the mechanical properties. The average storage moduli at 0.1 Hz and at 2 Hz (largest frequency before a drop-off was observed) were plotted against the average fluorescence intensity at each combination of genipin concentration and duration of incubation (Fig. 6). For both fluorescence measurement techniques and both frequencies, stiffness was correlated exponentially to intensity (Table I). The correlation coefficients were nearly identical for 0.1 and 2 Hz \( (R^2 = 0.808 \text{ and } 0.810, \text{ respectively, for measurements taken microscopically, and } R^2 = 0.782 \text{ and } 0.788, \text{ respectively for measurements taken spectrofluorometrically}) \). The exponential correlation curves shifted to the left slightly with increasing frequency, consistent with the increase in storage modulus. However, the resulting constants from the correlation were statistically indistinguishable.

FTIR characterization

The FTIR spectra of 10 mM genipin, “fully” genipin-crosslinked collagen (exposure to 10 mM genipin for 12 h and extensively rinsed of free genipin), and collagen during in situ crosslinking with 10 mM genipin are presented together in Figure 7. The spectrum of the genipin solution is dominated by three modes at 990, 1080, and 1635 cm\(^{-1}\), assigned to the ring C—H out-of-plane bend,\(^{32}\) ring C—H in-plane bend,\(^{32}\) and C=C double bond ring stretch modes\(^{32,33}\) of the core of the genipin molecule, respectively. The absorption at 1080 cm\(^{-1}\) may also include the C—O stretch mode of the primary alcohol on the genipin molecule.\(^{32}\) Additionally, the C—O—C asymmetric stretch and the CH\(_3\) bend of the methyl ester are observed at 1300 and 1443 cm\(^{-1}\), respectively. The 12 h crosslinked collagen spectrum features these modes, as well as bands at 1104 and 1370 cm\(^{-1}\) that are believed to be vibrational modes related to the formation of new bonds between genipin and the primary amines of lysine, hydroxylysine, or arginine residues in collagen. The band at 1370 cm\(^{-1}\) is assigned to the C—N stretch of the tertiary aromatic amine\(^{32,34}\) of the crosslinked genipin nitrogen iridoid\(^{35}\) that is bound covalently to the collagen. The broad, flat appearance of the crosslinking band at 1370 cm\(^{-1}\) in the 12-h spectrum is likely due to the flanking of two genipin molecule modes at 1360 and 1395 cm\(^{-1}\) (unassigned). The band at 1104 cm\(^{-1}\) is assigned to the C—N stretch of the tertiary nitrogen with the adjacent aliphatic carbon atom present in lysine or arginine residues.\(^{32,36}\) An absorption near 1104 cm\(^{-1}\) is also present in the unreacted genipin molecule as a shoulder to the absorption at 1080 cm\(^{-1}\). It is assigned to the vibrations of both the cyclic ether and secondary alcohol on the six-membered ring of the genipin molecule. When genipin reacts with collagen, both of these moieties are removed. Furthermore, the band at 1104 cm\(^{-1}\) in the 12 h crosslinked collagen is significantly stronger than the corresponding band in the spectrum of pure genipin (relative to the band at 1080 cm\(^{-1}\)) suggesting that this absorption band is mostly associated with modes formed as a result of crosslinking.
To better identify the origin of features present in the spectrum of 12 h crosslinked collagen, the changes in the collagen spectrum were monitored in situ during the first 4.5 h of crosslinking (Fig. 7). In this time-resolved experiment, spectral features were expected to increase due to: (1) diffusion of genipin into the region probed by the IR beam (the bottom surface of the collagen gel); and (2) crosslinking of collagen, leading to the appearance of new vibrational modes due to bonds formed during crosslinking. The in situ time-resolved spectra show the growth of several bands that are present in both crosslinked collagen and genipin, such as modes at 990, 1080, 1443, and 1633 cm\(^{-1}\). In addition, the beginning of the growth of a band centered near 1370 cm\(^{-1}\) is observed. This feature is only seen in crosslinked collagen. Figure 8 summarizes the time dependence of several absorbance features. Because of the proximity of the various genipin molecule and crosslinking bands, calculated band areas may include components of smaller bands adjacent to the dominant spectral feature. The 1080 cm\(^{-1}\) band area (spanning 1040–1180 cm\(^{-1}\)) likely includes the growth of a number of other smaller bands possibly including the crosslinking feature at 1104 cm\(^{-1}\), although it is too small to contribute substantially to band area. The feature at 1370 cm\(^{-1}\) is adjacent to genipin bands as stated earlier, and all are included in the area calculation (band complex spanning 1344–1414 cm\(^{-1}\)). To facilitate comparisons between

Figure 5. Fluorescence intensity (average ± std err) of genipin-crosslinked collagen measured using (A) epifluorescent microscopy (590 nm excitation, 630 nm emission) and (B) spectrofluorimetrically (590 nm excitation, 645 nm emission). For both, the intensity of fluorescence emission increased significantly with genipin concentration and duration of incubation (\(p < 0.001\)). Post hoc analysis (Fisher’s LSD test) revealed significant differences among all pairwise comparisons of concentrations except 5 mM versus 10 mM (max \(p = 0.199\)) and among all pairwise comparisons of duration except 4 h versus 6 h (\(p = 0.240\)).

Figure 6. Correlation of average storage modulus (± std err) with average fluorescence intensity (± std err) measured with epifluorescence microscopy (A) or spectrofluorimetrically (B) for 0.1 and 2 Hz. In all cases, strong, exponential correlations were observed, indicating that stiffness can be assayed optically following appropriate calibration. Increasing frequency shifted the correlation curve to the left.
trends in band growth, area absorbance values of the weaker band at 1370 cm\(^{-1}\) were scaled by a constant (indicated in the inset). Band area growth also differed in absolute value from run to run. Therefore, final \textit{in situ} values (\(t = 4.5\) h) of absorbance band areas amongst runs were normalized to a common value for each band, respectively. Similar increasing monotonic trends were observed for change in absorbance of genipin bands at 1080 and 1630 cm\(^{-1}\); however, the growth of the crosslinking band at 1370 cm\(^{-1}\) appeared to slow down within several hours. Indeed, the small area of the crosslinking band at 1370 cm\(^{-1}\) and its apparent slowing in growth are likely due to the relatively small number of genipin-to-collagen crosslinks in the gel that can form compared with the amount of genipin that diffuses to the region. A description of the modes marked in the spectra of Figure 7 is shown in Figure 9.

Cytotoxicity data

Most of the previous cytotoxicity studies of genipin had examined cell death following rinsing of genipin-crosslinked tissues or biomaterials prior to addition of cells. For tissue equivalent studies, knowledge of the cytotoxic effects of the genipin solution is required. Cytotoxicity studies using L929 fibroblasts indicated that genipin does cause significant cell death (ANOVA, \(p < 0.001\)) (Fig. 10). However, individual comparisons against the control condition (\textit{post hoc} analysis—Fisher’s LSD test) demonstrated that the adverse effects were limited to exposure of 5 mM (\(p < 0.001\)) and 10 mM (\(p < 0.001\)); though lower, cell numbers from samples incubated in 1 mM were statistically indistinguishable from controls (\(p = 0.26\)). The results at 12 h were consistent with the 24-h results.

<p>| Table I |
| Results of Storage Modulus—Fluorescence Intensity Correlations |</p>
<table>
<thead>
<tr>
<th>Fluorescence Measurement</th>
<th>Frequency (Hz)</th>
<th>(A) (±std err)</th>
<th>(B) (±std err)</th>
<th>(R^2)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Microscope</td>
<td>0.1</td>
<td>22.9 ± 7.33</td>
<td>3.05e-5 ± 4.30e-6</td>
<td>0.782</td>
</tr>
<tr>
<td>Microscope</td>
<td>2</td>
<td>24.3 ± 7.82</td>
<td>3.12e-5 ± 4.32e-6</td>
<td>0.788</td>
</tr>
<tr>
<td>Spectrofluorimeter</td>
<td>0.1</td>
<td>33.2 ± 8.36</td>
<td>8.34e-4 ± 1.1e-4</td>
<td>0.808</td>
</tr>
<tr>
<td>Spectrofluorimeter</td>
<td>2</td>
<td>35.7 ± 9.11</td>
<td>8.51e-4 ± 1.1e-4</td>
<td>0.810</td>
</tr>
</tbody>
</table>

\(G' = A \exp(B \times \text{Intensity})\)

Figure 7. IR Absorbance spectra of 10 mM genipin referenced to a spectrum of pure water (top); collagen after 12 h of crosslinking with 10 mM genipin and extensive rinsing, referenced to a spectrum of uncrosslinked collagen (second from top, scaled 2×) and collagen crosslinked with 10 mM genipin \textit{in situ}, 0.2, 0.4, 0.8, 1.7, 3.1, and 4.5 h after adding genipin, referenced to the initially genipin-free collagen gel (bottom). Several spectral features that are present in the genipin solution alone, and/or the crosslinked and rinsed collagen are seen to evolve during the \textit{in situ} crosslinking.
We have characterized the effects of genipin-induced crosslinking of collagen gels on rheological properties, fluorescence, spectroscopic changes, and cytotoxicity. Rheological measurements were performed at 1% shear strain amplitude over a range of shear rates, which is consistent with previous characterizations of collagen and other biopolymeric gels using similar techniques. We found that both the concentration and the duration of incubation in genipin significantly influenced the storage and loss moduli. The storage modulus measurements demonstrated a gradual increase with increasing frequency and were generally consistent with previous reports of collagen rheology. Loss modulus showed a gradual decrease with frequency. We further found that genipin-mediated crosslinking produced significant changes in fluorescence that are well-correlated to the stiffness, and that genipin has marked cytotoxic effects at concentrations of 5 mM and above. We conclude that genipin-induced crosslinking offers a simple alternative to improve the mechanical properties of tissue constructs, though caution must be taken to preserve cell viability.

We also observed a decline in the storage modulus (and increase in loss modulus) at larger frequencies that we associated with damage to the gel. In previous reports, where this trend was not observed, gels were generally prepared directly on the parallel plates. In our case, the lengthy incubations in genipin precluded this possibility, and instead gels were transferred to the rheometer. It is possible that the adhesion to the plates was not as optimal as when gels are directly prepared or that the transfer increased the potential for damage. Nonetheless, the observed decrease in storage modulus was consistent among samples and, interestingly, the “failure” properties of the gel improved with increasing concentration and crosslinking duration, indicating that it is a stress-based phenomenon.

Previously, genipin has been used to crosslink biological tissues, chitosan-based tissue equivalents, and gelatin with genipin concentrations ranging from 1 to 10 mM. In several of these studies, the influence of genipin-mediated crosslinking on cytotoxicity and/or cell viability has been evaluated for different cell types in several different conditions, each in the context of development of a genipin-crosslinked biomaterial, and relatively low toxicity has been identified following rinsing of the cross-linking.

Figure 8. Changes in absorbance band areas versus crosslinking time for some of the highlighted bands in Figure 7: two genipin bands (triangle, 1080 cm⁻¹ and circle, 1630 cm⁻¹), and a new genipin-to-collagen crosslinking feature (square, 1370 cm⁻¹). To allow all spectra to be viewed on a common plot, values of smaller band areas were scaled by a constant, as indicated in the inset. Crosslinking time began (at t = 0) when genipin reached the bottom of the collagen gel at the interface with silicon. Error bars combine two sources of errors: (1) baseline selection for area calculations, and (2) reproducibility of runs. End point area values of each band for different runs were normalized to a common value due to variation in absolute absorbance area possibly caused by variation in collagen density among samples.

Figure 9. Description of the modes marked in the spectra of Figure 7.
The rheological studies presented herein provide a screening of the effects of genipin on stiffness, but not a direct indication of the utility of the chemical for tissue engineering applications, particularly because we have not yet evaluated the effects of genipin on functional properties of cells. Additionally, the strains and rates used, while consistent with standard parallel plate rheology protocols, are insufficient to assess the utility of genipin for crosslinking bioartificial tissues that routinely experience finite deformations at accelerated rates. Future studies are aimed at identifying the appropriate timing and incubation durations to optimally influence the mechanical properties of cellular collagen constructs, including mechanical testing to larger strain levels at high rates more appropriate for tissue engineering applications. Previous characterizations with native tissues suggest that such treatments with genipin are plausible and may improve the mechanical properties of the constructs.17,19,20

Beyond providing significantly improved mechanical stiffness of acellular collagen gels, the ability to manipulate the mechanical properties of cellular tissue equivalents on a reasonably short time scale affords the opportunity to study, and potentially exploit, the phenotypic response of cells to changes in mechanical properties within a 3D tissue construct, if it is shown that genipin does not adversely influence cell viability and/or function. The past decade has shown increased focus on quantifying the behavior of cells grown on substrates or in systems of varying compliance, beginning with Pelham and Wang’s studies of fibroblast durotaxis using functionalized poly(acrylamide) gels.47 This system has been adapted to study neural cells,48 endothelial cells,49 and smooth muscle cells.50–52 In vivo, tissue cells reside within a three-dimensional (3D) matrix, which presents a significantly different set of environmental cues than when cells are cultured on a 2D substrate. Quantitative studies of the effects of mechanical properties on cell behavior in 3D, where cells are uniformly distributed throughout the tissue equivalent, rather than seeded on a gel and/or coaxed to invade the gel, have been limited to thin layers of collagen on top of poly(acrylamide) with controlled compliance to indirectly control the stiffness.53 Genipin can be used to crosslink the 3D tissue equivalents to influence the mechanical properties of the tissue matrix directly, allowing more complex shapes and boundary conditions to be investigated. For example, culturing cells within collagen gels seeded on poly(acrylamide) membranes naturally mimics a constrained system, where stress is generated at the poly(acrylamide)–collagen boundary as cells exert traction and attempt to contract the fibrillar network. However, it has been shown that the accumulation of network stress in a constrained system significantly affects the response of the resident cells.54 Genipin could be used to stiffen unconstrained, free-floating tissue equivalents as well as constrained ones to distinguish between the influence of the intrinsic mechanical properties of the fibrillar, extracellular matrix network and the mesostructural properties dictated by the network and its attachments/constraints to external entities.

Genipin also has the added novelty of producing crosslinks that appear blue and fluoresce allowing easy visualization and quantitation of crosslinking. The fluorigenic quality was first identified in forensics research that investigated genipin as a potential fin-

Figure 10. Cytotoxic effects of genipin. L929 fibroblasts entrapped in collagen gels were exposed for 24 h to culture medium with defined concentrations of genipin immediately upon completion of self-assembly. Live cells were labeled fluorescently with Calcein-AM, and the average number of cells (±std err) in a vertical field through the thickness of the gel was determined by manually counting the cells in a stack of several images taken through the height of the gel. Genipin was cytotoxic (ANOVA, \( p < 0.001 \)), but only a small fraction of cells were lost at 1 mM, and these results were not statistically different than the 0 mM control (Fisher’s LSD post hoc test, \( p = 0.26 \)).
gerprint reagent with increased sensitivity. We hypothesized that the same properties could be used to differentially indicate the degree of crosslinking in collagen gels, which would potentially enable the optical evaluation of mechanical properties. We measured the fluorescence intensity in parallel to mechanical properties in samples extensively rinsed to remove all free genipin, and found strong correlations between the genipin-generated fluorogenic properties and the mechanical properties of the collagen gels using both an epifluorescent microscope and a spectrofluorimetric plate reader. Spectrofluorimetric plate readers have advantages of high throughput and consistent measurement with no lag time between measurements of different samples. The plate reader will also capture the intensity signal through the thickness of the gel. However, only standard plate geometries can be used for measurement. Epifluorescent imaging is more cumbersome with each sample requiring manual focusing and measurement, but could prove useful for samples that do not fit into standard well-plate configurations, and for identifying any spatial variance in the degree of crosslinking. Traditional epifluorescent microscopy will also capture a signal through the thickness, but the signal will be strongest at the focal plane, while confocal microscopy could be used to pinpoint the fluorescence changes through the depth of the sample, as well.

The kinetics of the fluorescence changes roughly matched those in stiffness, but the intensity began to saturate at higher levels of crosslinking, leading to the exponential correlation of stiffness with fluorescence. In our experiments, fluorescence intensity was evaluated over a wide range of crosslinking regimens—from no crosslinking solution to incubation in 10 mM genipin for 12 h. To appropriately compare intensity values across this range, constant exposure settings were used for all conditions, and using the same settings necessary to measure low intensity levels at low levels of crosslinking can cause saturation at higher levels of crosslinking. A more sensitive calibration can easily be achieved by optimizing the exposure times for a narrower range of fluorescence.

The strong correlation of genipin-generated fluorescence to mechanical properties allows simple, noninvasive confirmation of mechanical properties for crosslinked gels/equivalents of various geometries that may not be particularly amenable to mechanical characterization to provide measures of within- and between-experiment variabilities. The ability to visualize crosslinks also enables direct observation of spatially varying crosslinking fields, such as defined patterns and gradients, and associated indirect assessment of the spatially varying mechanical properties. In all cases, the thickness of the actual sample and the ones used to generate a standard curve of fluorescence intensity versus crosslinking (duration or concentration) must be carefully considered, as a thicker sample (of the same collagen concentration) will generate increased fluorescence; the correlation presented is specifically derived from the samples probed in this study. A separate calibration is necessary for other sample sizes and/or collagen concentrations. Moreover, the introduction of cells and subsequent compaction of the collagen gel will alter the observed fluorescence by increasing fiber density. The fluorescent labeling of collagen via genipin also presents interesting opportunities to observe and measure collagen degradation via lost fluorescence. Thus, while the quantitation of stiffness via fluorescence may be best applied for prescribing and screening initial conditions to evoke specific, stiffness-driven behavior, the fluorogenic potential of genipin may also be used to evaluate matrix remodeling.

The colorimetric and fluorimetric properties of the crosslinked collagen are associated with molecular changes produced during crosslinking. In our study, the FTIR spectra include features at 1104 cm\(^{-1}\) (C—N stretch) and 1370 cm\(^{-1}\) (C—N stretch), that are neither characteristic of genipin nor collagen alone, and are, therefore, presumably associated with the crosslinked collagen and, perhaps, the color/fluorescence changes. The in situ FTIR demonstrated temporal changes in these features that paralleled the early changes in stiffness and fluorescence at 10 mM. However, due to limitations in the FTIR setup, the in situ spectroscopy could only be performed for \(~4.5\) h before evaporation began to introduce inconsistencies in the results, and a true correlation of stiffness-to-fluorescence-to-spectroscopy was not obtained. Interestingly, Tóyama et al. examined the intermediate pigment changes that occur upon reaction of genipin with methylamine. Brownish-red intermediates were associated with 2-methyl-4-carbomethoxy-2-pyrindine derivatives, which had a spectroscopic feature at 1630 cm\(^{-1}\). We also observed a peak at \(\sim1630\) cm\(^{-1}\) in the rinsed, crosslinked collagen, which is shifted slightly to the left of a corresponding peak at \(\sim1635\) cm\(^{-1}\) in the genipin solution. This shift may be artifactual due to water vibrational peak subtraction and/or due to double bonds, which contribute to the 1630/35 cm\(^{-1}\), being slightly affected due to their proximity to the covalent bonding upon crosslinking and subsequently causing a small 5 cm\(^{-1}\) shift for the average of all double bonds.

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References


