Influence of chondroitin sulfate and hyaluronic acid on structure, mechanical properties, and glioma invasion of collagen I gels

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1. Introduction

Extracellular matrix (ECM) is the complex environment composed of collagens, glycoproteins, glycosaminoglycans (GAGs) and proteoglycans (PGs) that surrounds and supports cells [1]. Type I collagen is the most common structural protein in mammalian tissue and has been widely used as a three dimensional (3D) ECM approximation in which to study cell behavior. Aside from the fact that collagen I environments are broadly physiologically relevant due to their dimensionality and the fact that they are composed of the most prevalent mammalian ECM component, collagen I environments also provide several other benefits for in vitro cell studies. First, collagen I forms fibrils that are easily visualizable with a variety of microscopies, such that when cells are embedded in these matrices, the cells, collagen fibrils, and their reciprocal interactions can be visualized simultaneously using multimodal microscopy [2–4]. Additionally, collagen I gels formed in vitro have tunable structure and mechanical properties controlled by gelation conditions including pH, temperature, and ionic strength [5–8]. Such tunability allows for construction of environments in which the independent effects of mechanical properties and network structure on cell behavior in 3D environments can be assessed [9–13].

While collagen I gels are generally used as single-component environments for in vitro studies of cell behavior, multi-component gels may more closely approximate the extracellular environment present in a particular tissue of interest. For example, while the migration and invasion of cells derived from high grade gliomas have been studied in collagen I environments [2,14–21], such environments are not especially good mimics of the brain tissue that gliomas invade in vivo. Unlike most tissue, brain does not contain large amounts of collagen I, instead being composed chiefly of neuronal and glial cells. However ECM does exist in the brain, both in the form of structural proteins and supporting GAGs and...
PGs, and some of the locations in which ECM is most prevalent in
brain tissue while retaining the benefits of collagen I environments,
we introduce CS or HA over a range of concentrations into collagen
solutions during fibrillogenesis to make collagen/CAG composites.
Such composites have been studied previously, though reports on
how CS and HA alter gel mechanical properties and network structure
are not in full agreement [28–38]. Because a subset of the
reported discrepancies may be related to differences in collagen
source, extraction technique, and preparation conditions, we
investigate the effects of CS and HA on collagen gels made from both
pepsin-treated (PT) and acid-solubilized (AS) collagen. PT collagen
lacks the telopeptides found in native collagen and is commonly
used in bioengineering applications due to its non-antigenicity. AS
collagen retains native telopeptides and interbrillar cross-links
and is more commonly used in biological studies. Differences
between networks composed of AS and PT collagen have begun to
attract attention [39,40], but potential differences between GAG/AS
collagen interactions and GAG/PT collagen interactions have not
been considered previously. We target a range of CS and HA
concentrations such that the GAG/collagen ratio spans from less
than one, consistent with the ratio in vasculature and other tissues,
to greater than ten, to approach the biochemical composition of
ECM surrounding gliomas and in normal brain parenchyma. Optical
microscopy is used to investigate potential changes to collagen fibril
structure and gel network structure that may occur in the presence
of the secondary ECM components, and oscillatory rheology is used
to monitor potential changes in gel viscoelasticity induced by the CS
or HA. We investigate whether correlations between gelation
kinetics, gel network structure, and gel storage modulus found in
pure collagen gels are also present in the GAG/collagen composites.
Glioma spheroid invasion in GAG/collagen gels at high GAG
concentrations is monitored. Performing these studies in both AS
and PT collagen helps isolate the biochemical effects of CS and HA on
glioma invasion.

2. Materials and methods

2.1. Materials

Acid-solubilized (AS) rat tail collagen I is obtained from BD Biosciences (San Jose, CA) and pepsin-treated (PT) bovine hide collagen I is obtained from Advanced BioMatrix (San Diego, CA). Hyaluronic acid sodium salt from streptococcus equi, chondroitin sulfate A sodium salt from bovine trachea, 10× Dulbecco’s Modified Eagle Media (DMEM), 7.5% v/v sodium bicarbonate, and 1 N sodium hydroxide are obtained from Sigma Aldrich (St. Louis, MO). 1 M Gibco HEPS buffer is obtained from Invitrogen (Carlsbad, CA). CS rat glioma cells are obtained from the American Type Culture Collection (Manassas, VA). The cell culture reagents, 1× DMEM (high glucose), heparin-inactivated fetal bovine serum (FBS), 100× antibiotic-antimycotic solution, and trypsin-EDTA, are obtained from Invitrogen.

2.2. Preparation of 3D collagen, collagen/HA, and collagen/CS gels

Pure collagen, collagen/HA, and collagen/CS gels are prepared at 4 °C. HA and CS are dissolved in water to prepare stock solutions of 10 mg/ml, 20 mg/ml, or 100 mg/ml (CS only). Appropriate volumes of collagen and HA stock solution or CS stock solution, depending on the final concentrations desired, are mixed with appropriate volumes of DMEM, sodium bicarbonate, HEPS buffer, deionized water, and sodium hydroxide to yield a pH of 7.4 and final percent volume concentrations of 0.05, 0.02 to 0.05, 2.5, and 2.5% DMEM, sodium bicarbonate, and HEPS, respectively. Such preparation conditions are used since they are supportive of cell culture. Solutions are well mixed and are placed in an incubator at 37 °C to gel or are held at 4 °C (for no more than 5 min after gel neutralization) before rheology, where they are gelled in situ. In 1.0 mg/ml collagen gels, final CS concentration ranges from 0.05 to 2.0% while final HA concentration ranges from 0.5 to 12.5 mg/ml, the highest HA concentration that could be achieved without leading to inhomogeneity in the gel.

2.3. Preparation of glioma spheroids

Cells are cultured at 37 °C with 5% carbon dioxide and are subcultured when 70–80% confluent. Multicellular tumor spheroids (MTSs) are formed by using the hanging drop procedure starting with cells diluted to 2.5–5.0 × 10^6 cells/ml. A 100 mm petri dish is filled with 10 ml of culture medium consisting of 1× DMEM (high glucose) containing 10% (v/v) FBS and a 1× (v/v) 100× antibiotic-antimycotic solution. Drops of 20 μl of the cell culture are placed on the lid of the Petri dish. The lid is inverted and incubated at room temperature for several hours. The drops are held in place by surface tension and the cells accumulate at the bottom of the droplet to form spheroids. The spheroids are then pipetted into collagen, collagen/HA, or collagen/CS solutions prepared as described above, after which they are placed in an incubator at 37 °C for gelation and maintenance of cell health. 50 μl of cell media is placed atop the spheroid-containing gels after each round of imaging (at 2, 24, 48, and 72 h after implantation).

2.4. Rheology

Rheological experiments are conducted on an AR-2000 rheometer with built-in temperature and gap calibration. A 1× acrylamide core geometry with a solvent trap is used. Time sweep experiments are conducted in oscillatory mode at a fixed frequency of 1 Hz. The collagen and collagen/GAG solutions are applied to the testing stage at 4 °C. The geometry is lowered, the solvent trap is added, and measurements begin when the temperature reaches 37 °C. Storage modulus (G’), frequency). For some PT collagen gels, to assure breakage of the developing gels during rheological measurements does not occur, gels are gelled on the rheometer for 45 min before oscillatory rheology measurements are performed. For all other gels, measuring moduli with either of these procedures leads to identical gels.

2.5. Microscopy and image analysis

Collagen or collagen/GAG solutions to be imaged are loaded just after neutralization of the gels. Gels containing collagen are imaged using a transmitted light microscope, and the solutions are brought to 37 °C and held there at least 1.25 h for AS collagen-based gels and at least 2 h for PT collagen-based gels, to assure gelation is complete. CRMs are recorded with an inverted confocal laser scanning microscope (Olympus Fluoview 300) with a 60× NA = 1.2 water objective. An argon ion laser at 488 nm is the excitation source, and the reflected light is detected on a photo-multiplier tube detector (PMT). Images per gel are collected and three gels at each concentration are investigated.

Mesh size is quantified from CRM images. In some images, a CRM artifact leading to a bright spot at the center of images, caused by a reflection in the optical path, is replaced with a copy of the upper left corner of the image in order to have representative fibers throughout the entirety of the image. The images are then thresholded in order to remove noise while retaining fiber pixels. The threshold is determined by comparing the intensities of the brightest background pixels to the dimmest fiber pixels through visual inspection. Pixels with intensity greater than threshold are labeled “on,” the distance between on pixels in each row and column is calculated, and the histogram of these distances is plotted. This distribution is fit to an exponential decay with the characteristic length scale determined by this fit. This size is defined as the mesh size. Mesh size determined in this manner has previously been shown to correlate well with another measure of mesh size [8].

Transmitted light microscopy images are used to determine the size of the spheroids, although this method is known to underestimate the size of the spheroids. However, this method is used here for comparison to the spheroids and a circle that circumscribes ~90% of the invasive cells, is determined for each image.
3. Results

3.1. Influence of CS and HA on network structure of collagen gels

CS or HA is added to both acid-solubilized (AS) and pepsin-treated (PT) collagen solutions to achieve final collagen content of 1.0 mg/ml and CS concentration ranging from 0 to 20 mg/ml or HA concentration ranging from 0 to 12.5 mg/ml. For collagen/HA solutions, homogeneous gels cannot be reliably prepared at HA concentrations higher than 12.5 mg/ml owing to the high viscosity of the HA. Solutions are then gelled at 37 °C.

Fig. 1 shows representative CRM images of 1.0 mg/ml AS and PT collagen gels as well as 1.0 mg/ml AS and PT collagen gels with 10 mg/ml CS or HA. Collagen fibril morphology in pure 1.0 mg/ml AS collagen gels is quite different from that in pure PT collagen gels. The PT gels are more homogeneous than are those of AS collagen, and the fibrils formed from PT collagen appear somewhat thicker, straighter, and less entangled than those of AS collagen (Fig. 1a,d). The average mesh size in the two types of gels is nearly identical (Fig. 2), as would be expected of gels of the same concentration so long as the same proportion of collagen is organized into visualized fibrils and those fibrils are of similar width in both preparations.

The network structure of AS collagen/CS gels as revealed by CRM imaging changes with CS concentration. The presence of CS at concentrations of as little as 0.5 mg/ml leads to imaged fibrils that appear straighter and brighter than those visualized in the pure collagen gels (data not shown). At CS concentrations of ≥5.0 mg/ml, imaged fibrillar structures appear significantly brighter and thicker than at lower CS concentrations and are present in some cases as bundled multi-fibrillar structures that we term fibers. Such fibers are similar to those present in pure AS gels formed at temperatures lower than 37 °C [8]. Assuming no change in proportion of collagen incorporated into fibrillar structures or average fibril length as a function of CS concentration, increased fibril thickness will lead to increased mesh size in the system. Such increased mesh size can be seen in Fig. 1b relative to Fig. 1a and is described quantitatively via mesh size analysis indicating an increase in mesh size from 5.1 to 12.0 μm for 1.0 mg/ml AS collagen with CS concentration from 0 to 20 mg/ml (Fig. 2).

Unlike in AS collagen, addition of CS to PT collagen causes little obvious change in fibril or network structure. Subtle increased brightness as a function of CS is apparent, suggesting a small increase in fibril thickness. For systems with a given amount of collagen present in fibrils, a small increase in average fibril thickness at fixed average fibril length will result in a small increase in mesh size, as is confirmed with mesh size analysis shown in Fig. 2.

Addition of HA to AS and PT collagen gels has markedly different effects on network structure than does addition of CS. For AS collagen, addition of HA results in an apparent increase in the number of fibrillar structures (Fig. 1c). The structures appear not only more numerous but also thinner and shorter than in the pure AS gels, and this is reflected quantitatively in the decreased mesh size as a function of HA concentration (Fig. 2). In PT collagen, addition of HA has subtler effects, though a small increase in fibril number and decreased mesh size is apparent (Fig. 1f and Fig. 2).

3.2. Influence of CS and HA on viscoelasticity of collagen gels

The storage ($G'$) and loss ($G''$) moduli of a viscoelastic material represent the energy stored in and dissipated by the material, respectively. These moduli can be measured using dynamic mechanical analysis (DMA) or oscillatory shear rheometry (OSR). The storage modulus ($G'$) is a measure of the elastic property of the material, while the loss modulus ($G''$) is a measure of the viscous property. The ratio of $G''/G'$, known as the loss tangent (tan δ), indicates the relative contribution of energy dissipation to the total energy stored in a material.

Fig. 1. CRM images of 1.0 mg/ml (a) AS collagen, (b) AS collagen with 10 mg/ml CS, (c) AS collagen with 10 mg/ml HA, (d) PT collagen, (e) PT collagen with 10 mg/ml CS, and (f) PT collagen with 10 mg/ml HA. Scale bars are 50 μm.
respectively. Fig. 3a presents change in equilibrium storage modulus for AS and PT collagen gels as a function of CS and HA concentration. Because pure AS and PT collagen gels have somewhat different storage moduli at 1.0 mg/ml (∼3 Pa for AS collagen and 6 Pa for PT collagen), normalized $G'$ is presented to highlight change relative to pure collagen. As a function of CS concentration, AS/CS collagen $G'$ values increase to a maximum at 5.0 mg/ml CS before decreasing to a minimum at the highest CS concentration employed, 20 mg/ml. For PT collagen gels, the changes as a function of CS concentration are similar to but of smaller magnitude than those seen for the AS gels. Just as the change in network structure of the gel with addition of CS is more apparent for AS collagen than PT collagen, so too is the change in storage modulus.

Conversely, even though AS collagen gel structure is more strongly affected by the presence of HA than is PT collagen gel structure, AS and PT collagen gels exhibit similar changes in equilibrium storage modulus with increasing amounts of HA (Fig. 3). Both AS and PT collagen gels with added HA exhibit $G'$ values that increase substantially with HA concentration. In addition to potential changes in collagen gel storage modulus as a function of secondary ECM concentration, the loss modulus may change as well. For AS and PT collagen systems with CS, there is little change in the loss tangent ($\tan \delta = G'/G''$) as a function of CS concentration (Fig. 3b). This indicates that the viscoelastic behavior of the composite collagen/CS gels is similar to that of pure collagen gels. In contrast, the loss tangent increases markedly upon adding HA to both AS and PT collagen (Fig. 3b). This occurs even though the absolute value of $G'$ increases substantially with added HA. The increase in the loss tangent is thus driven by a stark increase in loss modulus that occurs in these gels upon addition of HA. The increase in loss modulus is due primarily to viscous HA dispersed between the collagen fibrils. Indeed, pure 10 mg/ml HA, for example, has $G'$ of ∼5 Pa and $G''$ of ∼10 Pa (data not shown). As such, it appears that the viscoelastic response of collagen/HA gels at the concentrations investigated is dominated by the HA viscoelasticity while that of collagen/CS gels is dominated by collagen viscoelasticity.

In addition to changes in the equilibrium mechanical properties of the collagen gels, the presence of secondary ECM components may induce changes in gelation kinetics. For AS collagen, no clear change in gelation kinetics is seen upon the addition of CS. For pure AS collagen and AS/CS composites, the gels attain their equilibrium modulus within several minutes of the beginning of the rheological experiments in all cases, and the resolution of our experiments is insufficient to detect potential increases of gelation speed as driven by the CS. In PT collagen, which gels substantially more slowly than AS collagen, the time required for the storage and loss moduli to plateau decreases with the addition of even very small amounts of CS (Fig. 4). In these gels, the lag time (the time over which $G'$ is small and nearly constant), associated with nucleation, decreases with added CS. Additionally, the slope of the $G'$ rise, which correlates with growth and interconnection of fibrils, increases with
added CS. Thus, both the nucleation and growth phases of PT collagen gelation speed up upon addition of CS.

As for the addition of CS, the addition of HA does not noticeably affect the gelation speed of AS collagen, though this may again be related to the limited time resolution of the experiment. For PT collagen, in general, a decrease in gelation time as assessed by time to storage modulus plateau is seen; however, this is largely driven by an increase in initial $G'$ of the system rather than a decrease in lag time or increase of the slope of the growth phase. This again highlights the more complex nature of the viscoelasticity of the collagen/HA composites compared to the collagen/CS composites.

3.3. Correlation between structure and mechanical properties in collagen/CS gels

Previously, we have shown that both fibril or fiber diameter and mesh size are important in setting collagen gel storage modulus [8]. As such, if CS or HA alters either the fibrillar diameter or mesh size of the matrix, these changes alone are expected to affect the mechanical properties of the gel. In addition, the CS and HA themselves may alter the mechanical properties of the composite independently of their effect on collagen fibril and network structure. For cells embedded in collagen/CS or collagen/HA matrices, the potential effects of such changes on cell behavior must be considered before any conclusions can be drawn regarding the biochemical effects of the CS or HA on cell behavior.

Because HA itself is highly viscous under the conditions studied here while pure CS remains a Newtonian solution with low viscosity even at the highest concentrations employed, we consider whether changes in collagen/CS, but not collagen/HA, gel mechanical properties and structure are consistent with those found previously for pure collagen. First, we note that although CS clearly increases the rate of PT collagen gelation even at very low CS concentrations, this does not alter the structure of the collagen gel in the manner that increased gelation speed does in pure collagen gels. In pure collagen gels, increased gelation speed occurs with either increased collagen concentration or higher gelation temperature up to ~37 °C [5]. In both cases, fibril diameter decreases with increasing gelation speed [5,41]. For the PT collagen/CS gels, however, no obvious change in fibril diameter is apparent. Moreover, in AS collagen, where we cannot detect increased gelation speed as a function of CS due to limited time resolution, fibril bundling becomes apparent at high CS concentrations. For pure AS collagen, such bundling is only seen for low temperature gelation, where the gelation rate is significantly decreased from that at 37 °C [5,8]. For pure collagen gels, it is presumed that fast gelation correlates with thin fibrils because the growth portion of the nucleation and growth process of fibrillogenesis begins as soon as the nucleation centers reach critical size rather than allowing for larger nucleation centers to develop. In the collagen/CS gels, however, there is evidence for both increased gelation speed and fibril bundling. Thus, we propose that the CS alters AS collagen fibrils in a manner that allows for bundling through a mechanism distinct from that which allows for bundling in pure AS collagen gels at low gelation temperature.

We now assess whether the changes in equilibrium structure seen as a function of CS in the AS collagen/CS gels correlate with changes in mechanical properties in the same manner that changes in structure correlate with changes in mechanical properties in pure collagen gels. First, we note that for AS collagen with CS, $G'$ displays a maximum at the intermediate CS concentration of 5.0 mg/ml even though mesh size increases monotonically as CS increases from 0 to 20.0 mg/ml. This is similar to the behavior seen in AS collagen gels at a given concentration gelled at different temperatures, where mesh size increases monotonically as a function of decreasing gelation temperature but stiffness first increases and then decreases with lower gelation temperature. This behavior is attributed to the fact that network storage modulus depends on not only network mesh size but also the stiffness of the struts that comprise the network [8].

Qualitatively, the fiber bundles present in the AS collagen/CS gels with the highest CS concentration look similar to those present in pure AS gels constructed at 32 °C and do not exhibit as much bundling as those formed at lower temperatures [8]. We compare the correlation of mesh size and storage modulus for gels of 1.0 mg/ml AS collagen formed as a function of temperature and those formed at 37 °C with added CS (Fig. 5a). Figure 5a demonstrates that while the changes of mesh size and storage modulus are qualitatively similar for AS gels formed as a function of decreasing temperature and those formed in the presence of increasing CS, this behavior is distinct from that which allows for bundling in pure AS collagen gels at low gelation temperatures. This suggests that CS affects the mechanical properties of the fibrils or interfibrillar interactions beyond effects from the collagen bundling it induces. We now consider whether changes in collagen/CS, but not collagen/HA, gel mechanical properties and structure are consistent with those found previously for pure collagen. First, we note that although CS clearly increases the rate of PT collagen gelation even at very low CS concentrations, this does not alter the structure of the collagen gel in the manner that increased gelation speed does in pure collagen gels. In pure collagen gels, increased gelation speed occurs with either increased collagen concentration or higher gelation temperature up to ~37 °C [5]. In both cases, fibril diameter decreases with increasing gelation speed [5,41]. For the PT collagen/CS gels, however, no obvious change in fibril diameter is apparent. Moreover, in AS collagen, where we cannot detect increased gelation speed as a function of CS due to limited time resolution, fibril bundling becomes apparent at high CS concentrations. For pure AS collagen, such bundling is only seen for low temperature gelation, where the gelation rate is significantly decreased from that at 37 °C [5,8]. For pure collagen gels, it is presumed that fast gelation correlates with thin fibrils because the growth portion of the nucleation and growth process of fibrillogenesis begins as soon as the nucleation centers reach critical size rather than allowing for larger nucleation centers to develop. In the collagen/CS gels, however, there is evidence for both increased gelation speed and fibril bundling. Thus, we propose that the CS alters AS collagen fibrils in a manner that allows for bundling through a mechanism distinct from that which allows for bundling in pure AS collagen gels at low gelation temperature.

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of the fibril diameter present in the AS collagen/CS gels, as shown in Fig. 5b. The range of predicted fibril diameters is physically reasonable, with a prediction of 52 nm at 0 mg/ml CS (as compared to 56 nm measured for such gels using SEM previously) [8]) to a diameter of just over 90 nm, which correlates to bundles of ~4 fibrils. We conclude that AS collagen/CS gels exhibit changes in structure and network structural parameters that are consistent with those seen for pure AS collagen gels (Fig. 5b) in that both are well described by a semi-flexible polymer theory [42].

3.4. Influence of HA and CS on spheroid invasion in collagen matrices

As described above, incorporation of GAGs into collagen matrices can induce changes in structural, mechanical, and biochemical properties of the matrices, each of which can affect cell behavior. In order to elucidate the biochemical effects of HA and CS on glioma invasion, we study the growth of glioma spheroids in pure 1.0 mg/ml AS collagen and PT collagen gels as well as AS and PT collagen gels with 10.0 mg/ml CS or HA. Glioma MTSs are implanted in collagen and collagen/GAG gels. Two hours after implantation, collagen fibers surrounding the MTS are imaged (Fig. 6, bottom). The structure of the collagen networks surrounding the spheroids is similar to those found in cell-free gels (Fig. 1). MTS spheroid growth and invasion are monitored over time for 72 h. Invasive distance is calculated as the distance between the dense spheroid body and the average extent of invasive cells. Invasive character and distance in pure AS and PT collagen gels are similar, with significant invasion evident (Fig. 6a,b; Fig. 7). Cells demonstrate the migratory phenotype that has been seen previously for glioma cells in collagen gels, in which the cells extend along collagen fibers, exert traction on those fibers, release from the back end, and move forward [2]. In both AS and PT collagen gels, cells align collagen fibers into an astral pattern within several hours after spheroid placement in the collagen gel and then invade along these aligned fibers. While invasion in AS and PT gels is similar, invasion appears somewhat more aggressive in the AS gels, both in terms of invasive distance and invasive density, as the invasive front appears consistently denser in the AS gels than the PT gels of the same concentration.

Invasion in collagen/CS gels is inhibited relative to in pure AS and PT collagen gels, with fewer cells emerging from the dense spheroid body and their speed reduced relative to that in the pure collagen gels. While invasive distance is reduced in both AS/CS and PT/CS gels, the invasion in AS/CS matrices is more significantly hindered relative to its pure counterpart than is the invasion in PT/CS gels (Fig. 7). Indeed, despite qualitative differences between invasion in collagen and collagen/CS matrices, it is only differences in invasive distance between CS spheroids in AS and AS/CS gels that are statistically significant. Invasion in collagen/HA gels is qualitatively similar to that in pure collagen gels, though the cells adopt a somewhat less elongated morphology and the density of invading cells is somewhat higher than that in pure collagen gels. In PT collagen, the presence of HA somewhat reduces invasive distance relative to invasion in pure PT collagen, especially at 24 and 48 h. In AS collagen, this invasion inhibition is also seen at early times, but by 72 h after implantation, invasive distance is more substantial in AS/HA gels than in pure AS gels.

4. Discussion

Single-component collagen matrices have been widely used as 3D ECM mimics to study in vitro cell migration including glioma cell invasion [2,14–16,20,21]. However, gliomas in brain tissue encounter a variety of ECM components other than collagen as they invade either through brain parenchyma or along blood vessels, one of the preferential routes of invasion. GAGs including CS and HA are major components in brain ECM and are also found at elevated levels around brain tumors [43]. In order to better model the ECM of brain tissue for glioma invasion studies, CS and HA are incorporated into collagen gels.

**Fig. 5.** (a) Plateau storage modulus vs. mesh size for 1.0 mg/ml AS collagen gels with 0, 0.5, 1.0, 5.0, 10.0, and 20.0 mg/ml CS gelled at 37 °C (red circles) and for 1.0 mg/ml AS collagen gelled at 37 °C, 32 °C, 27 °C, or 22 °C (black squares). (b) Predicted fiber diameter of AS collagen as a function of CS concentration using the relationship between storage modulus, mesh size, and fibrillar diameter determined to fit behavior of pure AS collagen over a range of concentrations and gelation temperatures: log(G') = 0.871[log(ξ)+22/5]+9.4, with ξ mesh size and d fiber diameter [8] with d allowed to vary. Inset shows fit of AS collagen/CS data to the equation for the best-fit d values. Black squares are data from Reference [8] and red circles are data from AS collagen/CS gels (For interpretation of the references to color in this figure legend, the reader is referred to the web version of this article).
The structural and mechanical properties of collagen/CS or HA composite gels are studied over a range of concentrations that encompass physiological concentration of CS and HA in a variety of tissues including brain. We find that AS collagen networks are more significantly affected structurally and mechanically by the presence of CS than PT collagen networks. CS increases fiber thickness and mesh size substantially in AS collagen-based matrices. The bundling of fibrils into fibers that occurs at high CS concentrations in AS collagen but not in PT collagen is reminiscent of the behavior of these two types of collagen when gelled at temperatures of 32°C and below, where AS collagen fibrils bundle but PT collagen fibrils do not [8,41]. This suggests that fibril bundling is strongly inhibited by the absence of telopeptides even when gelation is performed at conditions that induce significant bundling in AS collagen. Our observations on gel structure in the presence of CS A are generally consistent with the structural changes found when small amounts of CS C are added to AS collagen I networks [33,35]. Our results differ somewhat from those of Douglas et al., where both CS A and CS C were found to decrease fibril diameter [32]. However, some fibril bundling in the presence of CS is evident in that study, and another study also suggests that high CS concentration can induce fibril bundling [28].

The storage modulus of PT collagen decreases slightly with CS concentration consistent with slightly increased mesh size in those gels. However, storage modulus of AS collagen shows a maximum at intermediate CS concentration in a manner consistent with the increase of mesh size and fiber thickness seen in these gels as well as in pure AS collagen (Fig. 5). This confirms that storage modulus is a function of both fiber dimension and organization [8] and that collagen/CS matrix mechanical properties are dominated by collagen viscoelasticity.

As with the addition of CS, AS collagen is more significantly affected structurally than is PT collagen upon addition of HA. More numerous and thinner collagen fibrils are apparent with increasing HA concentration in AS collagen and to a lesser extent in PT collagen. We note that while HA itself forms a meshwork, there is no evidence it forms structures that can be visualized with CRM; instead, as suggested previously, it is likely HA co-localizes with collagen fibrils and/or remains homogeneously dispersed in between fibrils [31,41,44]. In our previous work, it was found that introducing 1.2 mg/ml HA during PT collagen assembly slightly decreased fibril width and slightly enhanced storage modulus relative to pure PT collagen [41]. Here, at higher HA concentrations in PT collagen, we find no stronger effect on gel structure but a very strong effect on gel storage and viscous moduli, which are dominated by the HA mechanical response as reflected in the loss tangent. In AS collagen, the mechanical property changes induced by HA are similar to those seen in PT collagen, but the structural changes are more obvious, with more numerous fibrils present, resulting in decreased mesh size. Taking together the effects of CS and HA on AS and PT collagen, it is found that AS collagen is more substantially affected both structurally and mechanically by the presence of GAG molecules. This suggests that the non-helical ends of collagen may be crucial in controlling in vivo collagen self-assembly as it occurs in the presence of other ECM molecules.

We monitor the growth and invasion of spheroids in collagen gels with 10 mg/ml CS or HA because PT collagen gel structure is largely unaffected by the presence of the GAGs at that
concentration while AS collagen gel structure is substantially affected. Additionally, PT and AS collagen storage moduli are not strongly affected by the presence of CS at this concentration. Indeed, differences of stiffness much larger than that between pure collagen and collagen gels with 10 mg/ml CS were shown not to affect spheroid invasion so long as mesh size was held constant in a previous study in pure AS collagen gels [10]. For the addition of 10 mg/ml HA, both PT and AS collagen gel moduli are affected; however, the effect is due in large measure to the presence of the highly viscous HA rather than induced changes to the collagen network.

Given that PT gel topology and collagen network based viscoelasticity do not change significantly with added CS or HA, we interpret changes in invasive success of spheroids in PT collagen/GAG environments to be dominated by the biochemical changes induced by the presence of the GAGs. However, because AS collagen/GAG environments demonstrate significant structural change in the presence of 10 mg/ml CS and HA, differences in growth in these environments is expected to reflect both the biochemical presence of the GAGs as well as potential effects of the changes in the underlying network structure.

Our results indicate that C6 spheroids invade somewhat more aggressively in terms of invasive speed and cell density in AS collagen gels than in PT collagen gels. CS inhibits glioma invasion in both AS and PT collagen, though the difference is only statistically significant in the AS/CS gels. In PT collagen, any inhibition of invasion induced by the presence of CS is assumed to be related to CS biochemistry rather than changes to the structure and/or mechanical properties of collagen gels induced by CS, since structural and mechanical properties in PT collagen change very little with addition of CS. In AS collagen, there is a more substantial inhibition of invasion, potentially reflecting the biochemical effect of the CS plus the effect of a larger mesh size environment composed of fibril bundles, factors which may independently inhibit the migration of glioma cells through limiting cell ability to exert traction on and remodel the surrounding collagen structures [10]. The mechanism by which the biochemistry of CS may reduce invasion is not immediately clear. It has been found that CS A can increase adhesion of osteoblasts to collagen fibrils [32]. While such interactions do not necessarily suggest decreased invasion, if the presence of CS enhances cell-collagen adhesion substantially, this may prevent cell release from the collagen fibrils, thus preventing invasion deep into the collagen matrix [45,46].

While CS inhibits glioma invasion, HA appears to have limited effect on invasion in PT and AS collagen gels, with some evidence that HA encourages growth at long times in AS collagen gels. It has been shown previously that HA can facilitate glioma invasion [23,47,48] though one study shows HA oligomers may limit the malignant behavior of gliomas [49]. In vivo, it has been suggested that HA can facilitate invasion through absorbing water and enlarging the pericellular space, providing space through which the cells could migrate. In vitro in 1.0 mg/ml gels, which display a mesh size amenable to cell migration, this factor is not expected to come into play. Indeed, we see a decrease in mesh size as a function of increasing HA concentration in both PT and AS collagen-based matrices. The fact that HA appears to slow invasion at early times in both PT and AS based matrices may suggest that the increased viscoelasticity of the system at early times inhibits invasion. Late invasion, however, accelerates in both types of gels, with a more noticeable increase occurring in AS collagen-based gels. These effects may be due to a combination of HA’s effects on the biochemistry of the environment and a contribution from the increased number and decreased width of fibrils seen in the presence of HA, as in this mesh size range increased fibril number enhances invasive density and speed by allowing for more cells to generate traction on and move along particular fibrillar structures. This is consistent with the higher density of invasive cells seen in the HA containing gels relative to either the pure collagen gels or the CS containing gels.

5. Conclusion

Composite matrices containing collagen and either CS or HA were constructed to enhance similarity between the in vitro environment and brain tissue ECM. The structure and mechanical properties of collagen/CS and collagen/HA gels were studied via CRM and rheology. It was found that AS collagen network structure is affected by both CS and HA, while PT collagen network structure is largely unaffected. Moreover, the mechanical properties of AS collagen-based gels are more substantially affected by the presence of CS than are PT gels, suggesting the source of the change is the alteration in collagen network structure. Indeed, the relationship between microstructure and storage modulus in AS/CS gels demonstrates that, in cases in which the secondary component does not itself have a complex viscoelastic behavior, gel storage modulus is determined by both fiber diameter and network mesh size. The effects of CS and HA on glioma invasion were then studied in collagen/GAG matrices with structure both similar to (PT collagen-based gels) and different from (AS collagen-based gels) those of pure collagen matrices. From these studies, it appears that cell invasion in AS collagen-based gels is more substantially affected by the presence of secondary ECM components than in PT collagen-based gels. This highlights the fact that not only the biochemical effect but also any changes in collagen network structure induced by the secondary ECM component must be considered when evaluating cell behavior in composite environments.

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