Molecular Origin of the Elastic State of Aqueous Hyaluronic Acid

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ABSTRACT: The macroscopic mechanical properties of biological hydrogels are broadly studied and successfully mimicked in synthetic materials, but little is known about the molecular interactions that mediate these properties. Here, we use two-dimensional infrared spectroscopy to study the pH-induced gelation of hyaluronic acid, a ubiquitous biopolymer, which undergoes a transition from a viscous to an elastic state in a narrow pH range around 2.5. We find that the gelation originates from the enhanced formation of strong interchain connections, consisting of a double amide–COOH hydrogen bond and an N–D–COO− hydrogen bond on the adjacent sugars of the hyaluronan disaccharide unit. We confirm the enhanced interchain connectivity in the elastic state by atomic force microscopy imaging.

INTRODUCTION

Hydrogels regulate the mechanical properties of cells and tissues and thereby play an important role in many biological and pathophysiological processes, ranging from stem cell differentiation and the degradation of aging cartilage to wound healing. An important feature of natural hydrogels is their ability to tune their viscoelastic properties in response to environmental stimuli, such as the solution pH, temperature, and salt conditions. Up to now, it has been difficult to connect the macroscopic response of hydrogels to the underlying molecular interactions, e.g., hydrogen-bond and hydrophobic interactions, thus limiting the capability to predict and tune the properties of hydrogels.

One of the few techniques able to track interactions at the molecular scale is two-dimensional infrared spectroscopy (2D-IR). 2D-IR shares with two-dimensional nuclear magnetic resonance the ability to resolve molecular couplings and dynamics, but 2D-IR can resolve these interactions at a much shorter time scale. Molecular interactions such as hydrogen bonds or electrostatic forces usually change on a time scale of picoseconds, and with 2D-IR, snapshots of these interactions can be taken with femtosecond time resolution.

Here, we use 2D-IR to identify the molecular interactions that drive the pH-triggered gelation of hyaluronic acid (HA), a natural polyelectrolyte that has received widespread attention, being one of the main components of the extracellular matrix. It is important for cell–cell communication and also plays an important role in many biological processes, such as joint lubrication, skin hydration, and cell migration, whereas its dysregulation contributes to cancer development. Due to its biocompatibility, hyaluronic acid is also widely used as a scaffold for cell culture and drug delivery. Hyaluronic acid is a paradigmatic example of a responsive natural hydrogel: it displays a sharp transition in mechanical behavior at pH 2.5, switching from a viscous state to an elastic gel, denoted as the "putty state." The mechanism underlying this remarkable liquid-to-gel transition has been investigated for over almost a century with macroscopic techniques that map the dependence of the viscoelastic properties on pH, temperature, and ionic strength. Based on these studies, it has been proposed that at pH = 2.5, the net charge of the polyelectrolyte chains may be sufficiently suppressed to enable the formation of interchain hydrogen-bond interactions that would act as crosslinks. This principle was suggested to be applicable to the wider class of polysaccharides, but there has been no direct evidence for this putative pH-triggered hydrogen-bonding mechanism.

EXPERIMENTAL SECTION

Sample Preparation. The samples were prepared in a glass vial by adding water (or heavy water), NaCl (0.15 M), and HCl (or DCl) to achieve different molarities ranging from 0 to 80 mM of acid. All of these chemicals were provided by Sigma Aldrich. A specific amount of hyaluronic acid sodium salt in powder form from Streptococcus equi bacteria from Sigma Aldrich (1.5–1.8 > 1 MDa), or from Lifecore

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Biomedical \((M_w \sim 150 \text{ kDa})\), was added to the solution to reach a final concentration ranging from 10 to 20 mg/mL. The samples were left to equilibrate at room temperature for at least 24 h before measuring. Samples were stored for at most 1 week at a temperature of 4 °C, to slow down the hydrolytic degradation process.  

**Rheology.** Rheology measurements were performed with a stress-controlled rheometer (Anton Paar MCR 501), equipped with stainless steel parallel plates with a diameter of 40 mm. The experiments were performed at a gap size of 100 μm and at a temperature of 22 °C set by a Peltier system. Samples left to equilibrate for 1 week to reach homogeneity were loaded on the plate using a spatula. After thermal equilibration, the elastic and viscous shear moduli were probed by performing oscillatory shear measurements at an oscillation frequency of 0.5 Hz and a strain amplitude of 0.5%, which is well within the linear viscoelastic regime for hyaluronan. The reported results are averages of at least three measurements for each pH. For each sample, the pH was measured using a pH meter (Hanna Instruments).

**Fourier-Transform Infrared (FTIR).** All linear absorption measurements were performed using a Bruker Vertex 80v FTIR spectrometer, equipped with a liquid-nitrogen-cooled mercury cadmium telluride (MCT) detector. The spectra were recorded under a nitrogen atmosphere at a wavelength resolution of 3 cm⁻¹. For every spectrum, 100 scans were averaged. In all measurements, a standard sample cell with a path length of 100 or 75 μm was used. The reported spectra were corrected for the absorption of the solvent background.  

**2D-IR.** In 2D-IR experiments, we performed one-color experiments by pumping and probing around 1680 cm⁻¹ and two-color experiments by pumping at 1680 cm⁻¹ and probing at 1450 cm⁻¹. The home-built setup that we use has been described before.  

**RESULTS**

**Rheology and Infrared Spectroscopy.** Hyaluronic acid has a relatively simple linear structure and homogeneous composition with repeating disaccharides composed of N-acetyl-glucosamine and glucuronic acid (illustrated in the inset for each condition tested.

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**Figure 1.** Hyaluronic acid solutions undergo a sol–gel transition in a narrow pH range. (a) A tube inversion assay shows that hyaluronic acid solutions prepared at pH 1.6 exhibit viscous flow (s), whereas solutions prepared at pH 2.5 form a viscoelastic gel (p) that flows only on time scales beyond 2 min. (b) The pH dependence of the viscous and elastic modulus of hyaluronic acid in heavy water solutions. We observe a sharp peak in the elastic shear modulus at pH 2.5. The inset shows that hyaluronic acid is a polymer of disaccharides, themselves composed of N-glucuronic acid and N-acetyl-D-glucosamine. (c) Linear infrared spectra for hyaluronic acid solutions in D₂O at pH values ranging between 1.6 and 7. The infrared spectrum at pH = 2.5 is represented by the thick solid line. Between 1550 and 1760 cm⁻¹, we observe three bands: the antisymmetric stretching mode of the carboxylate anion group, \(\nu_{\text{AM.I, COOD}}\), at 1607 cm⁻¹, the amide I vibration, \(\nu_{\text{AM.I}}\), at 1633 cm⁻¹, and the carboxylic acid stretching mode, \(\nu_{\text{COOH}}\), at 1726 cm⁻¹. The inset shows the fractions of COO⁻ and COOD groups determined from the IR spectra as a function of pH. Thick lines represent the expected fractions based on acid–base equilibrium equations (Supporting Information), assuming a pKₐ = 2.9. We find good agreement between the measured and expected carboxylic and carbonate fractions, with about 25% deprotonation of the carboxyl groups at pH = 2.5.
of Figure 1b). The pH-dependent viscoelastic behavior of hyaluronic acid solutions stands out clearly in tube inversion assays, as shown in Figure 1a. Whereas the sample prepared in a solution state (“s”) at pH 1.6 readily flows, the sample prepared in the putty state (“p”) at pH 2.5 only flows on a time scale of minutes. This large difference in time scales originates from a pH-triggered transition in rheological properties from liquid to gel. To study the pH dependence of this transition in quantitative detail, we performed rheological experiments on semidilute solutions of high and low molecular weight hyaluronic acid. We studied solutions in heavy water (D$_2$O), since heavy water is used as a solvent in 2D-IR studies of the amide and carbonyl vibrations. 2D-IR cannot be applied to solutions in H$_2$O, as the amide and carbonyl vibrations of hyaluronic acid absorb in the same infrared frequency region as the bending vibration of H$_2$O (between 1600 and 1700 cm$^{-1}$). In all experiments, the pD values are converted to pH values (see Supporting Information Appendix, Figure S1b), which shows that the elastic and the viscous moduli sharply increase in a narrow range around a pH of 2.5, revealing a liquid–gel transition. This transition occurs at a pH value just below the pK$_w$ which is about 2.9. We find a qualitatively similar pH dependence of the viscous and elastic moduli for hyaluronan solutions in H$_2$O (see Supporting Information Appendix, Figure S2), which shows that the isotope composition of the solvent has a negligible effect on the mechanical properties.

We studied the protonation state of hyaluronan as a function of pH with linear IR spectroscopy (Figure 1c). The relative intensities of the carboxylic acid and carboxylate anion bands of hyaluronan directly reflect the protonation state of hyaluronan. The antisymmetric vibration of the carboxylate anion, $\nu$$_{\text{at,COO}}$, absorbs at 1607 cm$^{-1}$, and the amide I band, $\nu$$_{\text{AM.I}}$, absorbs at 1633 cm$^{-1}$. The C=O vibration of the COOD group, which will be denoted as $\nu$$_{\text{COOD}}$, absorbs around 1726 cm$^{-1}$. At pH = 2.5, the COO$^-$ and COOD groups are present in a ratio of 1:4 (inset in Figure 1c), which corresponds to a pK$_w$ of 2.9, as reported in the literature.

Two-Dimensional Infrared Spectroscopy. We use 2D-IR spectroscopy to study the intermolecular interactions among hyaluronic acid chains that lead to the solution–gel transition. We excite the amide and the carbonyl vibrations with a strong femtosecond infrared pulse pair (~100 fs, 4 $\mu$J per pulse). This excitation induces transient absorption changes that are probed with a weaker (0.35 $\mu$J) single femtosecond probing pulse that is delayed by a time $T_m$. The excitation and probe pulses are centered at 1680 cm$^{-1}$ with a bandwidth of 200 cm$^{-1}$. The excitation and probe spectra thus have sufficient spectral width to cover the $\nu = 0$ to $\nu = 1$ transition (bleach) and the $\nu = 1$ to $\nu = 2$ transition (excited state absorption, or esa) of the $\nu$$_{\text{at,COO}}$, $\nu$$_{\text{COOD}}$, and $\nu$$_{\text{AM.I}}$ vibrations.

Figure 2a,b compare the 2D-IR spectra measured for the solution (pH = 1.6) and the putty state (pH = 2.5), both for $T_m$ = 1.5 ps. Bleaching (increased transmission) is indicated in blue, whereas increased absorption (esa) is indicated in red. The bleach of the $\nu = 0$ to $\nu = 1$ transition and the esa of $\nu$$_{\text{AM.I}}$ are centered at 1639 and 1619 cm$^{-1}$, respectively, along the probe frequency axis, whereas for $\nu$$_{\text{COOD}}$, these signals are centered at 1731 and 1703 cm$^{-1}$, respectively (see also Supporting Information Appendix, Table S2). The bleaching and esa of the $\nu$$_{\text{at,COO}}^{-1}$ mode are in the low-frequency wing of the bleach and esa of the amide I vibration, and lead to an elongation of the bleaching and the induced absorption signals to the lower left corner of the 2D-IR spectrum. We observe additional signals in the off-diagonal regions of the 2D-IR spectrum, indicating that the carboxyl and amide modes are coupled. In the upper pump-frequency region, the cross-peak signal represents the effect of the excitation of the $\nu$$_{\text{COOD}}$ vibration on the spectrum of the $\nu$$_{\text{AM.I}}$ mode. In the lower pump-frequency region, the spectral shape of the carboxylic acid vibration is elongated along the pump axis (Figure S7). This effect reduces the visibility of the upward cross-peak, which represents the effect of the excitation of the $\nu$$_{\text{AM.I}}$ vibration on the spectrum of the $\nu$$_{\text{COOD}}$ vibration. For this reason, we focus our analysis on the downward cross-peak that represents the effect of the excitation of the carboxylic acid vibration on the $\nu$$_{\text{AM.I}}$ vibration. The excitation of the $\nu$$_{\text{COOD}}$ vibration also leads to a cross-peak signal with the amide II mode ($\nu$$_{\text{AM.II}}$) at 1490 cm$^{-1}$. This signal results from the excitation of low-frequency modes following the relaxation of the $\nu$$_{\text{COOD}}$ mode. The excitation of these low-frequency modes affects the absorption spectra of both the $\nu$$_{\text{COOD}}$ and $\nu$$_{\text{AM.II}}$ modes (see SI Appendix, Figure S6a,b).

To learn more about the mechanism that underlies the cross-peak signals of the amide I modes following excitation of the COOD vibration, we compare the dependence on time delay for the solution and the putty states (Figure 3a). We normalized the transients to the bleaching signal of the carboxylic acid vibration at early time delay (~0.2 ps). From ~1.5 ps on, the decay of the transients is quite similar to that of the $\nu$$_{\text{COOD}}$= $\nu$$_{\text{AM.I}}$ cross-peak, showing a time constant of ~3
Figure 3. a) Decay traces of the cross-peak in the $\nu_{AM.I}$ vibrational region for hyaluronic acid at a concentration of 20 mg/mL in the solution and putty states. The decay traces are normalized to the carboxylic acid intensity at early time delay. The solid and dashed black lines are fits to the relaxation model (described in the Supporting Information), for the putty and solution states, respectively. The blue lines represent the fast component: thick and dashed lines correspond, respectively, to the putty and solution states. The initial amplitude of the fast component (indicated with dashed lines) is nearly two times larger in the putty state ($c_{\text{ff,putty}} = 0.13 \pm 0.01$) than in the solution state ($c_{\text{ff,sol}} = 0.077 \pm 0.005$). The red line represents the contribution of the slow component, whose initial amplitude $c_{\text{ff}}$ is similar for the putty and solution states ($c_{\text{ff,putty}} = c_{\text{ff,sol}} = 0.024 \pm 0.004$). Values reported are the mean and standard deviation obtained by averaging over three different independent measurements. (b) Schematic of the relaxation model describing the cross-peak dynamics. Red and blue lines refer, respectively, to the slow component (due to thermalization) and fast component (due to energy transfer) of the cross-peak dynamics. The value shown is the average with standard deviation for six independent measurements.

ps. However, the transients of the $\nu_{COOD}$$-\nu_{AM.I}$ cross-peak signals have their maximum at 0.3–0.5 ps, a much earlier delay time than the one observed for the $\nu_{COOD}$$-\nu_{AM.II}$ cross-peak ($\sim$1–1.5 ps). In addition, the transients of the $\nu_{COOD}$$-\nu_{AM.I}$ cross-peak show an additional faster decay component with a time constant of $\sim$0.6 ps. Interestingly, the transient of the $\nu_{COOD}$$-\nu_{AM.I}$ cross-peaks has a much higher intensity at early delay times for the putty state than for the solution state. This observation, together with the dynamics observed for the $\nu_{COOD}$$-\nu_{AM.II}$ cross-peak, indicates that the $\nu_{COOD}$$-\nu_{AM.I}$ cross-peak signal contains contributions from two distinct mechanisms: (1) energy transfer from the excited COOD vibration to the amide I vibration; (2) anharmonic coupling to low-frequency modes that affect the vibrational spectrum of the amide I mode. The relaxation model corresponding to these two mechanisms is illustrated in Figure 3b.

The cross-peak signal arising from the energy transfer rises with a time constant $T_{\text{ent}}$ representing the time scale of energy transfer from the excited COOD vibration to the accepting amide I vibration, and decays with the vibrational lifetime, $T_{1AM.I}$ of the amide I vibration. This direct energy transfer does not occur for the amide II mode because there is a rather large energy difference between the $\nu_{COOD}$ and $\nu_{AM.II}$ vibrations. We extract the value of $T_{1AM.I}$ from the diagonal signal of the amide I mode in the 2D-IR spectrum. We find that $T_{1AM.I}$ is comparable to the vibrational lifetime of the COOD vibration: $T_{1AM.I} = 0.65 \pm 0.1$ ps. The time constants of the second mechanism involving anharmonic coupling to low-frequency modes can be extracted from a fit to the transient of the $\nu_{COOD}$$-\nu_{AM.I}$ cross-peak (SI Appendix, Figure S6b). We thus fit the transients of the $\nu_{COOD}$$-\nu_{AM.I}$ cross-peak to the following expression containing the lifetimes of the COOD vibration, $T_{1COOD}$, the amide I vibration, $T_{1AM.I}$, and the low-frequency-modes, $T_{1LFM}$.

$$
\frac{\Delta t_{\text{cp}}(t)}{\Delta t_{\text{COOD}}(t)} = e_{\text{anh}} \left(\frac{T_{1LFM}}{T_{1COOD} - T_{1LFM}}\right) \left(e^{-t/T_{1COOD}} - e^{-t/T_{1LFM}}\right) + e_{\text{ent}} \left(\frac{T_{1AM.I}}{T_{\text{ent}} - T_{1AM.I}}\right) \left(e^{-t/T_{\text{ent}}} - e^{-t/T_{1AM.I}}\right)
$$

Here, $\Delta t_{\text{cp}}(t)/\Delta t_{\text{COOD}}(t)$ is the intensity of the cross-peak normalized to the intensity of the $\nu_{COOD}$ vibration at early time delay, $t = 200$ fs. The parameters $e_{\text{anh}}$ and $e_{\text{ent}}$ represent the cross-peak amplitudes associated with the anharmonic coupling and direct energy transfer mechanisms, respectively. $T_{\text{ent}}$ and $e_{\text{ent}}$ are the only free parameters. As shown in Figure 3a, $e_{\text{ent}}$ is nearly 2-fold larger for the putty state than for the solution state, whereas $e_{\text{anh}}$ is comparable for the two states. For the time constant $T_{\text{ent}}$ we obtain a value of 150 ± 50 fs. The energy transfer to the amide I mode constitutes an additional channel for vibrational relaxation of the COOD mode. The fact that we do not observe a significant difference in $T_{1COOD}$ between the putty and the solution state indicates that this energy transfer process occurs only for a minor fraction of the COOD groups. From the values of $e_{\text{ent}}$ we estimate this fraction to be 13 ± 2% for the putty state and 7 ± 2% for the solution state. The time constant $T_{\text{ent}}$ also does not change upon the formation of the putty state, which shows that the efficiency of the energy transfer remains the same. Hence, upon formation of the putty state only the number of COOD groups showing energy transfer increases.

The ultrafast energy transfer between the C=O vibration of the COOD group and the amide I vibration of the amide group can be well explained from the presence of a hydrogen bond between the COOD and the amide groups. To get more information on the nature of this hydrogen bond, we compared the intensity of the $\nu_{COOD}$$-\nu_{AM.I}$ cross-peak signal for polarizations of the probing pulse parallel and perpendicular to the polarization of the excitation pulse. The ratio of these two signals reveals the relative orientation of the transition dipole moments of the excited and probed vibrations that form the cross-peak signal. From the ratio of the signals at $T_{\text{ent}}$ between 0.3 and 1 ps (where the energy transfer mechanism is predominant), we find that the angle between the transition dipoles of $\nu_{COOD}$ and $\nu_{AM.I}$ is $\sim$20° (SI Appendix, Figure S9). As the transition dipole moment of $\nu_{AM.I}$ is at an angle of 20° relative to the C=O bond of the amide group, we conclude that the hydrogen-bond configuration involves a parallel (or anti-parallel) alignment between the C=O groups of the carboxylic acid and amide, forming two closely spaced intermolecular hydrogen bonds between the amide and carboxylic acid groups, i.e., C=O\cdots O=C–OD\cdots O=C–ND and N\cdots O=C–OD\cdots O=C–OD. This geometry, in which the amide and carboxylic acid groups act both as the hydrogen-bond donor and acceptor stabilizing the hydrogen-bonded...
structure formed by these two molecular groups, constitutes a strong interchain connection. We conclude that the putty state at pH = 2.5 contains a 2-fold higher density of these strong double hydrogen bonds than the solution state at pH = 1.6.

To study the role of the COO− groups in the formation of the putty state, we also measure 2D-IR spectra where we excite and probe the ν\text{\textsubscript{\text{as,COO}}} and the ν\text{\textsubscript{AM.I}} vibrations. Since the absorption bands of these two vibrations overlap, we subtract the parallel signal from three times the perpendicular signal. In the resulting difference spectrum, the diagonal peaks are strongly suppressed since the corresponding parallel signal is ~3 times larger than the perpendicular signal. This procedure is thus ideally suited to reveal the presence of cross-peak signals of vibrations that are differently oriented. This strategy has been extensively used in 2D-IR experiments on proteins to resolve vibrational couplings. Figure 4a reports the resulting difference 2D-IR spectrum at pH = 2.5 (putty state). On the diagonal, we observe some residual signal due to orientational relaxation of the COO− and amide molecular groups. In addition, we also see clear off-diagonal signals (indicated by the yellow rectangles). In Figure 4b, we plot the transient absorption spectrum at pH = 2.5 (black circles) obtained from Figure 4a by averaging the signals with pump frequencies between 1600 and 1620 cm\textsuperscript{-1}, corresponding to the excitation of the ν\text{\textsubscript{\text{as,COO}}} vibration. The resulting spectrum is compared with the transient absorption spectrum at pH = 2.9 that is obtained using the same procedure (SI Appendix, Figure S12). The transient absorption spectrum at pH 2.5 clearly shows an enhanced absorption signal around the amide I vibrational frequency, which implies that the excitation of the ν\text{\textsubscript{\text{as,COO}}} vibration yields an enhanced response of the ν\text{\textsubscript{AM.I}} vibration at pH = 2.5. This means that in the putty state (pH = 2.5), the interaction between the COO− and amide molecular groups is enhanced. A similar enhancement is observed for the other cross-peak signal, i.e., exciting the amide I vibration and probing the COO− mode (SI Appendix, Figure S11f). Hence, our results show that the formation of single N–D–COO− hydrogen bonds is enhanced in the putty state, in addition to the enhanced presence of double amide–COOH hydrogen bonds in this state.

**Atomic Force Spectroscopy (AFM).** We also perform atomic force microscopy (AFM) imaging of hyaluronic acid solutions at different pH values to investigate to what extent the additional hydrogen bonds in the putty state are formed within or between hyaluronic acid chains (Figure 5a,b). At pH = 7, we mainly observe pearl-necklace conformations: thin chains alternating with thicker globular configurations. The globular portions are likely due to the low affinity for mica. At pH = 2.5, the AFM images clearly show a much larger density of thick filaments than at pH = 7, indicative of interchain interactions. To quantify the extent of lateral association of hyaluronan polymers, we measured the heights of the filaments at pH = 7 and 2.5 (Figure 5c). At neutral pH, we find a Gaussian height distribution centered at an average filament height $h_0 = 0.37$ nm. This value is consistent with the expected thickness of a single hyaluronan chain. At pH = 2.5, we find a much broader distribution, and $h_0 = 0.53$ nm, meaning that the filaments are thicker. Since the length of the filaments at pH = 2.5 is not changed compared with that at pH = 7, we conclude that this thickening is mainly due to enhanced interchain interactions that laterally associate the strands.

**DISCUSSION**

The two-dimensional infrared (2D-IR) spectroscopy measurements show that the remarkable pH-triggered gelation of hyaluronic acid at pH 2.5 involves the enhanced formation of double hydrogen bonds between carboxylic acid and amide groups, and of strong single hydrogen bonds between a carboxylate anion (which is a stronger hydrogen-bond acceptor) and amide groups. The enhanced formation of hydrogen bonds is intimately related and even enabled by changes in the structure and electrostatic interactions of the hyaluronan polymer chains.

An important question is why the gelation occurs only in a narrow pH range. In particular, one may wonder why the
gelation does not occur for a larger pH range below 2.5, as at lower pH, there will be even more carboxylic acid groups and thus potentially even more strong double hydrogen bonds with the amide groups located on a different chain.

The answer to this question can be found in the special structure of the hyaluronic acid disaccharide unit, which contains a carboxylic acid group and an amide group on neighboring sugar units (see the inset of Figure 1b). The formation of a double amide–carboxylic acid hydrogen-bond structure involves a quite strict local positioning of the hyaluronic acid chains, which prevents the formation of a second double hydrogen-bond structure at the adjacent saccharide units. Hence, the following mechanistic picture arises (Figure 6): at pH = 2.5, the partial deprotonation of the carboxylic acid groups allows for a high probability of forming a strong double amide–COOH hydrogen bond (green box) and an adjacent strong N–D–COO⁻ hydrogen bond (orange box), together forming a very strong interchain connection. In the single N–D–COO⁻ hydrogen bond, the N–D of an amide group acts as the hydrogen bond donor and the COO⁻ anion group as the hydrogen-bond acceptor. For a sufficient length and concentration of the hyaluronic acid chains, this enhanced hydrogen-bond formation will yield a long-range connected network that behaves as an elastic solid, the putty state. This state is not formed at lower pH, because at these pH values, very few carboxylic acid groups are deprotonated and the strong double amide–COOH hydrogen bond can only be combined with a relatively weak single N–D–COOH or single COOH–amide hydrogen bond on the neighboring sugar units, thus forming a weaker interchain connection. As a result, fewer interchain connections are formed (ca. 2-fold less at pH ≈ 1.6), and the overall connectivity is too low to form an elastic gel. Also, at pH values above 2.5, the putty state is not formed because a large fraction of the carboxylic acid groups is deprotonated, which implies that no double amide–COOH hydrogen bonds can be formed. In addition, Coulomb repulsion will prevent the formation of N–D⋯O–C hydrogen bonds on neighboring sugar units. As a result, the density of interchain connections is again too low to allow for gel formation.

![Figure 6](image)

**Figure 6.** Schematic images showing the transition from a weakly hydrogen-bonded network of hyaluronan at pH < 2.5 (top left) to a strongly hydrogen-bonded network at pH = 2.5 (right), to isolated single chains at pH > 2.5 (bottom left). At pH = 2.5, the balance between hydrogen-bond and Coulomb repulsion interactions is optimal, and allows the formation of hydrogen bonds between the carboxylic acid and amide groups observed as the fast time component in the cross-peak dynamics in 2D-IR. Green shaded squares indicate the formation of hydrogen bonds between amide and protonated carboxyl groups, whereas orange shaded squares depict the formation of hydrogen bonds between amide and deprotonated carboxyl groups.

**CONCLUSIONS**

In summary, we studied the molecular origin of the putty state of the biologically important polysaccharide hyaluronic acid, which undergoes a sol–gel transformation only in a narrow range around pH = 2.5. Using two-dimensional infrared (2D-IR) spectroscopy, we find that this remarkable transition is accompanied by the enhanced formation of strong interchain connections that consist of a strong double amide–COOH hydrogen bond (green box in Figure 6) and a strong N–D–COO⁻ hydrogen bond (orange box in Figure 6) on the adjacent sugar groups of the hyaluronan disaccharide unit. These strong collective interchain connections can only form when the carboxylic acid groups are partially deprotonated, which explains why the putty state is observed only in a narrow pH range. The enhanced interchain connectivity in the putty state is confirmed by AFM measurements that reveal the association of hyaluronan into thick strands in this state.

This study illustrates that the combination of 2D-IR spectroscopy, AFM, and rheology constitutes a unique tool to identify the molecular mechanisms by which hydrogels respond to environmental stimuli. This combined multiscale approach has great potential to investigate the molecular origin of the biological functionalities of other natural polyelectrolytes and to assist in the rational design of supramolecular biomaterials with life-like functionalities.

**ASSOCIATED CONTENT**

**Supporting Information**

The Supporting Information is available free of charge on the ACS Publications website at DOI: 10.1021/acs.jpcb.9b00982.

Data analysis methods; hyaluronic acid gel pictures; rheology experiments in water; linear infrared spectra; anisotropy data; 2D-IR spectra; AFM measurements (PDF)

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**Notes**

The authors declare no competing financial interest.
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