Hyaluronan orders water molecules in its nanoscale extended hydration shells

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Hyaluronan (HA) is an anionic, highly hydrated bio-polyelectrolyte found in the extracellular environment, like the synovial fluid between joints. We explore the extended hydration shell structure of HA in water using femtosecond elastic second-harmonic scattering (fs-ESHS). HA enhances orientational water-water correlations. Angle-resolved fs-ESHS measurements and nonlinear optical modeling show that HA behaves like a flexible chain surrounded by extended shells of orientationally correlated water. We describe several ways to determine the concentration-dependent size and shape of a polyelectrolyte in water, using the amount of water oriented by the polyelectrolyte charges as a contrast agent. The spatial extent of the hydration shell is determined via temperature-dependent measurements and can reach up to 475 nm, corresponding to a length of 1600 water molecules. A strong isotope effect, stemming from nuclear quantum effects, is observed when light water (H₂O) is replaced by heavy water (D₂O), amounting to a factor of 4.3 in the scattered SH intensity.

INTRODUCTION

Water has a key function in biology as it is the primary solvent of life. It actively participates in biochemical transformations such as protein folding, molecular recognition, self-assembly, and joint lubrication (1–4). To elucidate the role that various constituents of the body play in sustaining life, it is important to take into account their interaction with the aqueous environment. One example where water plays a very important role is in the extracellular matrix of epithelial and connective tissues. The matrix is rich in hyaluronan (HA), an anionic poly-disaccharide composed of β-N-acetylglucosamine and β-glucuronic acid (displayed in Fig. 1A) (5). HA has the ability to retain up to 10× its dry weight of water (6–11) and creates a gel-like pericellular coat (12, 13). HA is a component in nearly every part of the body, varying in concentration from 10 ng/ml (~12 pM) to ~3 mg/ml (3.5 μM) (5), and its hydration is thought to be key for the functioning of moving joints (12, 14, 15). HA has been studied in solution with linear light (16) and neutron scattering (16, 17) at concentrations above 100 μg/ml, with mid-sized (tens to thousands of kilodaltons) macromolecules. At concentrations <100 μg/ml (~<0.1 μM), the size and shape of the polyelectrolyte are unknown. Several theoretical models (18–20) exist for predicting the shape of polyelectrolytes, taking into account both thermal motion and electrostatics; however, these are only partially valid because they require rigid macromolecules (such as DNA), a condition that is not met for HA. Calorimetric (21, 22), vibrational spectroscopic (23–25), and compressibility (26) studies indicate slower hydrogen (H)–bonding dynamics and different thermodynamic properties of the hydrating water compared to bulk water. However, the observed effects of HA are limited to the first or second hydration shell. Recent developments in nonlinear optical spectroscopy allow direct probing of the orientational order in the H-bonding network of water over nanoscale distances and under extreme dilution (27). This enables further investigation of HA hydration under more dilute conditions, exploring weaker and longer-ranged interactions.

The structure of water near ions and charged groups is perturbed relative to neat water due to well-known ion–dipole and H–bonding interactions (28–30). However, the spatial extent over which ions perturb the structure of water is still a subject of debate (27, 31–33). Ions are traditionally assumed to have a short-range ordering effect, influencing the structure of, at most, the first three layers of hydrating water. This notion is supported by dielectric and terahertz spectroscopy (34, 35), x-ray (36) and neutron scattering (37), vibrational spectroscopies (38–40), and computational simulations (41, 42). However, recent studies based on femtosecond elastic second-harmonic scattering (fs-ESHS) experiments (27, 43, 44), supported by computational studies (45, 46), are changing this long-held notion that ion–water interactions are limited to several angstroms. Unlike most experimental techniques, which are often inherently limited to subnanometer length scales or equivalently require >100 mM solute concentrations, fs-ESHS directly probes spatial correlations over length scales comparable to the wavelength of the light used and in very dilute solutions (47). Using fs-ESHS, a nonspecific long-range (~20 nm) influence of simple electrolyte ions on the hydrogen (H)–bonding network of water molecules was found that starts at ionic strengths as low as 10 μM (27). This observation was explained as an electric field–induced perturbation in the water–water orientational correlations. Moreover, there is a noticeable nuclear quantum effect that is yet to be explained.

Here, we measure the orientational order of water in HA solutions and explore the extended hydration shell structure of HA in light and heavy water using fs-ESHS. We observe that water–water orientational correlations are enhanced by HA substantially more than by simple salts. The increased order in water is measurable from HA concentrations ranging from <1 pM up to ~1 μM, with an excess ionic strength range from 1 nM to 10 mM. A substantial isotope effect is observed when light water (H₂O) is replaced by heavy water (D₂O). The onset concentration increases in D₂O by a factor of 2.4, and the scattered intensity is 4.3× lower than in H₂O. Angle-resolved fs-ESHS measurements were performed to investigate the structure and size of water–HA complexes. Using nonlinear light scattering theory to model the intensity generated by the oriented
water, two different spherical radii are determined, $R$ and $R'$. $R$ is the size of an on average spherical object where the electrostatic potential has a maximum. At $R'$, this electrostatic potential has decayed to 2% of its maximum value. Temperature-dependent angle-resolved fs-ESHS measurements were performed to determine the size of the extended hydration shell at the concentration of maximum fs-ESHS intensity. The extended hydration shell ($R_h$) is determined by the water that is associated directly with the HA chain by molecular interactions, and $R < R_h < R'$. The extended spherical hydration shell $R_h$ that includes the HA macromolecule spans up to 475 nm in $H_2O$ and 260 nm in $D_2O$, corresponding to a line of 1600 (928) water molecules in $H_2O$ ($D_2O$).

RESULTS AND DISCUSSION

fs-ESHS from HA in $H_2O$ and $D_2O$

To provide context, we first recount a recent finding concerning fs-ESHS measurements performed on aqueous solutions of simple electrolytes. Figure 1B shows the fs-ESHS intensity recorded at a scattering angle $\theta$ of 90° for NaCl in $H_2O$ (red triangles) and $D_2O$ (tilted blue squares). The data were adapted from (27, 48). The increase in the fs-ESHS intensity was attributed to arise from an increase in the orientational order in the H-bonding network of water due to the interaction of the ionic electrostatic field with the H-bonded network. The intensity starts increasing from a salt concentration of $\sim$10 $\mu$M and saturates at $\sim$1 mM. Electrolytes dissolved in $D_2O$ show the same trend as in $H_2O$ but with a $\sim$6x higher onset concentration and a 36% smaller saturation value, relative to the pure $H_2O$ or $D_2O$ solvent. This slight restructuring of the H-bonding network also manifests itself on the macroscopic length scale as the Jones-Ray effect (32), an anomalous decrease of the surface tension at the fs-ESHS saturation concentrations. The increase in orientational order in the bulk solution gives rise to an entropic penalty, which reduces the surface tension (27, 32, 49).

Figure 1C shows fs-ESHS data obtained in the same manner for HA (molecular weight, 1145 kDa) dissolved in $H_2O$ (red circles) and $D_2O$ (blue squares) plotted as a function of calculated excess ionic strength and HA concentration in nM. fs-ESHS is a nonresonant process in which each molecular group contributes a comparable amount to the total nonlinear polarization in the solution. The intensity that is measured scales with the absolute square of the nonlinear polarization. At an HA concentration of 11.6 nM (13.3 $\mu$g/ml, $I_{excess} = 6.3 \mu$M) for $H_2O$ and $I_{PPP(\text{HA})/PPP, \text{water}} = 1.45$ at a $c_{\text{HA}} = 27$ nM (32 $\mu$g/ml, $I_{excess} = 39.5 \mu$M) for $D_2O$.

Note that these HA concentrations are far below those probed previously with other techniques (>100 $\mu$g/ml, and 117 nM only in $H_2O$; (16, 17, 20) and summarized in table S1).

The increased magnitude of the fs-ESHS intensity shows that the amount of orientational order in the water as induced by HA is much greater for HA compared to NaCl. This indicates that the
interactions of the electrostatic field from the ionic groups on the HA polymers with the H-bonding network in water is much stronger than the same type of interaction in the electrolyte solution. This is caused by the spatial arrangement of the ionic groups on the polymer: Each HA monomer, 1.02 nm in size, contains one negative charge (Fig. 1A) so that HA has a linear charge density of ~1 e⁻/nm with the entire chain containing >3000 units. As the spacing of charges is larger than the Bjerrum length of water (0.71 nm), the ionic groups of HA are ionized. The combination of electrostatic and thermal interactions significantly complicates the description of HA polyelectrolyte structure, and although several models exist (16, 18–20), the structure of HA in solution is unknown in the concentration range of our investigation. It is clear, however, that unlike ions of simple salts, which are statistically distributed in solution, the charges on HA cannot move freely as they are fixed by the polymer chain that links them. These charges are therefore spatially correlated over a certain distance due to the covalent bonding between individual monomers on a single polymer chain and the limited degrees of freedom that the chain has due to bending, thermal motion. In addition, intrachain electrostatic repulsion further limits the flexibility and spatial positioning of the HA chains (19).

The concentration of such a large number of electric charges in a small region of space generates much stronger localized electrostatic fields than statistically distributed free ions. This leads to a much stronger effect of the electrostatic field on the H-bonding network of water and thus a much larger fs-ESHS response of HA compared to NaCl.

**Nuclear quantum effects**

As can be seen in Fig. 1C, the fs-ESHS response of HA in H₂O and D₂O is remarkably different. The fact that replacing light with heavy water in HA solutions significantly alters the fs-ESHS response indicates that the HA chains do not simply order water molecules via a charge-dipole interaction: Charge-dipole interactions would not be detectably affected by a change in the isotope of the water as both water isotopes have nearly identical dielectric constants and dipole moments (50). The isotope effect in the fs-ESHS response points to a more complex process in which the two primary and orthogonal degrees of freedom to break and form hydrogen bonds, H-bond stretching and H-bond rotation (or libration), are decoupled. The magnitude of the difference further points toward long-range interactions involving many water molecules. In addition, this nuclear quantum effect involves the electric field of HA interacting with the total H-bonding network of water, changing the orientational water-water correlations in the solution (27, 48). Compared to simple electrolytes, HA induces these effects at much lower concentrations [10 μM versus ~10 pM of HA (I_{excess} = 10^{-8} M for H₂O) and 100 μM versus ~1 nM of HA (I_{excess} = 10^{-6} for D₂O)] and with a much higher magnitude (1.05 versus 6.4 for H₂O and 1.0 versus 1.4 for D₂O, at the same ionic strength). Because a similar effect coincides with a small but significant change in a macroscopic property of the solution for simple salts, the much bigger change here might lead to much bigger changes in macroscopic properties of hydrated polyelectrolytes. One such candidate is the viscosity of the liquid (51), which depends on the reorientation of water, and has been correlated to the coherent fs-ESHS response of concentrated aqueous electrolyte solutions (52). Although it is very clear from the experiment what the parameters for describing this nuclear quantum effect should be, up until now, theoretical modeling has not yet been able to capture this behavior (27, 51). To describe the phenomenon observed in Fig. 2 (B and C), it will be necessary to describe HA and its extended surrounding water molecules on a quantum chemistry level. Although these computations have been done on a single water molecule embedded in an environment that can be described by a classical force field (53), the need to do this for a much bigger cluster involving HA and many waters would involve computations on a scale that is currently not feasible. Therefore, in what follows, we revert to classical nonlinear light scattering theory (54, 55) to model the structure of HA polyelectrolyte hydration in H₂O. We will, therefore, describe angle-resolved and temperature-dependent second-harmonic scattering data of HA in H₂O and extract relevant length scales that we will then compare with what is known about the conformation of HA (18, 19), thereby adding structural information of HA and its hydration in a range that is currently inaccessible by other methods.

**Fig. 2. Scattering patterns of HA solutions are concentration dependent.** (A) Angle-resolved fs-ESHS measurements of HA solutions (molecular weight, 1349 kDa) in H₂O measured at three concentrations showing a strong concentration dependence of the angular distribution of the scattered SH light. More concentrated HA solutions scatter at higher angles. 0° corresponds to a 4.8 mM NaCl solution. The excess ionic strengths of the other solutions are 0.4 μM (0.3 μg/ml), 3.8 μM (3 μg/ml), and 12.5 μM (10 μg/ml). More patterns are shown in the Supplementary Materials (fig. S1). (B) Integrated fs-ESHS intensity as a function of HA concentration. The solid curve shows the predicted behavior based on the proposed model of fs-ESHS and is calculated using Eq. 3. (C) Angle of maximum fs-ESHS intensity plotted as a function of HA concentration. The scattering angle at maximum intensity (θ_{max}) shifts to higher angles with increasing concentration, which is equivalent to scattering from smaller objects. The solid lines were calculated using Eq. 4. The parameters used for the modeling are as follows: R = 5 μm⁻¹ and I₀ = 5 μm (see Materials and Methods). The experimental data were measured in the PPP polarization combination. The error bar is representative for all data points.
Angle-resolved fs-ESHS

To characterize the flexible structure of hydrated HA, we measured angle-resolved fs-ESHS in the entire forward half-plane (from \( \theta = -90^\circ \) to \( \theta = +90^\circ \)) for HA dissolved in H_2O. In contrast to single-angle measurements, the angle-resolved measurements provide information about the shape of the emitted scattered light pattern, which is important for further analysis. Three representative angle-resolved scattering patterns of HA solutions at concentrations of 0.2, 2.2, and 7.4 nM (0.3, 3, and 10 \( \mu \)g/ml, with an excess ionic strength of 0.4, 3.8, and 12.5 mM) and one pattern of a NaCl solution (4.8 mM) are shown in Fig. 2A. Patterns for all of the measured HA concentrations are shown in fig. S3, showing NaCl patterns (fig. S3A) and HA patterns (fig. S3B). The patterns were recorded with all beams polarized along the horizontal scattering plane (PPP polarization combination). The shape of the scattering pattern of a NaCl solution does not vary with salt concentration, as can be seen in fig. S3A. In contrast, the shape and intensity of the HA patterns substantially change with increasing HA concentration. The variations in the data of Fig. 2A can be captured by two key parameters: The integrated fs-ESHS intensity \( S_{int} \), calculated by integrating the scattering patterns in the forward half-plane, is shown in Fig. 2B. The angle of maximum scattering intensity \( \theta_{max} \) is shown in Fig. 2C. Both metrics are illustrated in more detail in fig. S3C. \( S_{int} \) increases with HA concentration up to ~5 \( \mu \)g/ml (6.25 \( \mu \)M) and then drops and levels off, very similar to the data measured at \( \theta = 90^\circ \) (Fig. 1B). \( \theta_{max} \) shifts monotonously from \( \theta_{max}= 18^\circ \) to \( \theta_{max}= 65^\circ \) with increasing HA concentration. To understand the unusual concentration dependence, we model the two essential properties of the scattering by using nonlinear light scattering theory. The solid lines in Fig. 2 (B and C) correspond to the result of this model, which we describe next.

Modeling of the fs-ESHS scattering patterns

Nonlinear light scattering theory has been developed for a variety of systems, including SHS from an isotropic distribution of molecules (56), used to describe the incoherent hyper-Rayleigh scattering from liquids (27, 44, 52), and coherent SHS from spherical surfaces or infinitely thin shells around isotropic particles in an isotropic solution (57–60). HA molecules in solution are macromolecular and have a size range well over a nanometer so that hydrated HA can be thought of as an on-average soft particle. Our measurements report on many such macromolecules in the laser beam focus (with a beam waist of 35 \( \mu \)m, \(~4.7 \times 10^5\) HA molecules at 11.6 nM) in combination with \(~2.2 \times 10^{11}\) water molecules. The nonresonant SH polarizability reports on every noncentrosymmetric molecule with an equal magnitude (61), and therefore, the SH intensity reports on the water. This means that the coherent water signal determines the nonlinear optical response of our experiment. Furthermore, the averaged water response is influenced by the electrostatic field on the water orientation. It is therefore possible to approach this modeling either by using a model that treats water as a dipolar liquid (27), performs averaging of the hyperpolarizability tensors \( \beta^{(3)} \) and \( \beta^{(3)} \), and removes the incoherent part or by directly using the coherent part through the susceptibility tensors \( \chi^{(3)} \) and \( \chi^{(3)} \) already available from the nonlinear light scattering theory of spherical objects in solution (47). Although the choice is eventually arbitrary, we use the latter one, applying the Rayleigh-Gans-Debye (RGD) approximation, in which the refractive index contrast between a particle with radius \( R \) and the medium is small (62). We will first describe the model as it is known and then adjust it to describe the current system. The coherent response is characterized by a second-order surface susceptibility \( \chi_{SS}^{(3)} \), which captures interfacial chemical interactions. Later models have included the effect of a nonzero surface potential \( \Phi_0 \) and ionic strength (via \( \kappa^{-1} \)) (63). The electrostatic potential decays to ~2% of its surface value at \( R \) over a distance of 4\( \kappa^{-1} \); thus, \( R = R + 4 \kappa^{-1} \) (54). This introduces an effective third-order susceptibility \( \chi^{(3)} \), which takes into account the contribution to the coherent SH response of water molecules that are oriented by an electrostatic field. For the present nonresonant excitation, \( \chi^{(3)} \) has only a single value, \( \chi^{(3)} = 10.3 \times 10^{-22} \text{ m}^2/\text{V}^2 \) (64). Figure 3A shows an illustration of the relevant size parameters in this model. The dimensions \( R \) and \( R' \) are determined by the electrostatic potential \( \Phi \); \( R \) is the radial distance from where the electrostatic potential \( \Phi \) starts to decay into the solution, and \( R' \) is the radial distance where this potential has decayed to 2% of its maximum value, the surface potential, \( \Phi_0 \). For SHS from solid particles, droplets, and liposomes, \( R \) is comparable to the linear light scattering or hydrodynamic radius (62, 64–67). The reason for this is that the refractive index and the electrostatic potential experience discontinuities at approximately the same radial distance. Section S3 contains a more detailed description of RGD SHS from a sphere, which results in the following type of expression for the scattered SH response \( S(\theta) \) in terms of the intensity scattered by a particle dispersion in PPP polarization \( I(\theta)_{PPP} \), disp, normalized by the intensity scattered by pure water in the SSS polarization combination \( I(\theta)_{SSS,H_2O} \).
The single-particle response for noninteracting particles, determined by the form factor \( F(q, R, \kappa, \chi_{\text{SSS}}^{(3)}, \Phi_0) \), is multiplied by the particle density \( N_p \) of the scattering wave vector \( q \) defined as the difference between the wave vectors of the scattered SH wave vector and the sum of the two fundamental wave vectors. This leads to a characteristic two-lobe pattern, similar to those plotted in Fig. 2A (54-58, 62, 65). Within this model, the position of the lobes (represented by \( \theta_{\text{max}} \)) shifts to higher angles with decreasing particle radius. For example, taking a wavelength of 1000 nm, and \( \chi_{\text{SSS}}^{(3)} = 1 \), results in \( \theta_{\text{max}} > 50^\circ \) for \( R = 50 \) nm, while \( \theta_{\text{max}} < 20^\circ \) is reached when \( R = 500 \) nm.

Having described this model, we first consider its applicability to polyelectrolytes in solution and then describe the implemented size models, such as the Odijk-Skolnick-Fixman (OSF) theory (18-19). We will treat the proportionality constant \( \alpha \) as a fitting parameter to obtain \( R \). Such a choice is in accordance with other size models, such as the Odijk-Skolnick-Fixman (OSF) theory (18, 19), although the exponent on \( \kappa^{-1} \) varies depending on the polyelectrolyte rigidity, the concentration regime, and the ionic strength (70, 71). We will discuss the general issue of polyelectrolyte size and the difference between \( R \) and the radius of gyration, as well as the difference in contrast mechanism further in the “Polyelectrolyte size: Comparing linear and nonlinear scattering” section.

To compare the model (Eq. 2) with the experimental data in Fig. 2 (B and C), we calculate the total fs-ESHS signal by numerically integrating the scattering pattern in the forward half-plane

\[
S_{\text{int}}(\theta) = \int_{-\pi/2}^{\pi/2} S(\theta) \ d\theta
\]

The maximum scattering angle \( \theta_{\text{max}} \) of Fig. 2C can be obtained by computing the maximum of the scattering pattern in the forward half-plane, i.e., by differentiating and numerically solving for \( \theta \).
The peak integrated intensity at different HA chains are suppressed as well, reducing the structure factor. Second, the extent of the hydration relaxation is weakened (reducing \( R_s \)). Third, long-range charge-charge correlations between different HA chains are suppressed as well, reducing the structure factor. The peak integrated intensity at \( c_{HA} = 3.71 \) nM (5 \( \mu \)g/ml) is reached when the positive contribution from an increasing number density of the number of HA chains and their extended hydration shells. At the same time, increasing the HA concentration adds counter-ions to the solution that increases the excess ionic hydration shells. At the same time, increasing the HA concentration (in micrograms per milliliter). It can be seen that the radius of gyration \( r_g \) through a common theory, one would be able to connect both types of contrast mechanism. However, the most widely used model, OSF theory \( (18, 19) \), describes the polyelectrolyte size with a combination of thermal and electrostatic interactions. OSF theory is valid when \( a \ll \lambda_B L_0 \) \( (20) \), where \( a \) is the separation between the charges \( (a=1.02 \text{ nm for HA}), \lambda_B \) is the Bjerrum length (\( \lambda_B=0.71 \text{ nm} \)), and \( L_0 \) is the intrinsic persistence length of HA \( (L_0=8\text{ nm} \; (17)) \). For HA, \( \sqrt[3]{\lambda_B L_0} = 2.4 \text{ nm} \), meaning that the chain is assumed to be stiffer than it actually is. In addition, the relationship between a computed value such as the end-to-end distance \( r_{ee} \) and the experimentally measured radius of gyration is not well determined \( (72) \): For a neutral flexible chain, the ratio between \( r_{ee} \) and \( r_g \) is \( \sqrt{6} \), whereas for a stiff rod it is \( \sqrt{12} \). Here, we take \( \sqrt{9} = 3 \) as determined by \( (72) \), who found this value to be true for short HA chains (<64 monomers). Although there are ~3000 monomers per chain in this study, we assume that the number of monomers does not influence this relation significantly.

Figure 4A presents a comparison of the variation of \( R \) (i.e., the size of the polymer as determined by the radial distance from which the electrostatic potential decays into the solution) and \( R' \) (i.e., the size of the polymer chain and the region with a nonnegligible electrostatic potential), together with the radius of gyration as determined by OSF theory from the end-to-end distance \( (17-19) \). The contour length \( \lambda_c \) (i.e., the length at maximum extension) and the free radius per HA, \( R_{free} \), are also plotted. \( R_{free} \) is calculated as follows: \( R_{free} = \left( \frac{M_{HA}}{4\pi c_{HA} N_A} \right)^{1/3} \), where \( M_{HA} \) is the average molar weight of HA \( (M_{HA}=1349\text{ kDa}) \), \( N_A \) is Avogadro’s constant, and \( c_{HA} \) is the HA concentration (in micrograms per milliliter). It can be seen that the contour length is larger than all other size parameters, where \( \lambda_c \) represents the end-to-end distance of a fully stretched chain. The computed \( r_g \) is larger than both \( R \) and \( R' \) for the entire concentration

\[
\frac{dS(\theta)}{d\theta} \bigg|_{\theta_{\text{max}}} = 0
\]  

(4)

The computed \( S_{int} \) and \( \theta_{\text{max}} \) values are shown in Fig. 2 (B and C) as solid lines. Using the single fit parameter \( \alpha \), which takes a best-fit value of \( \alpha = 5 \), the curve for the total fs-ESHS intensity produced by this model (Fig. 2B) matches well with the measured values (circles), as well as the maximum scattering angle as a function of HA concentration (Fig. 2C). Considering that both the integrated intensity and the angle of maximum intensity are found with the same single fit parameter \( \alpha \) suggests that our model captures the essentials of the fs-ESHS response.

**Concentration dependence**

The modeling provides insight into the unusual concentration dependence of the fs-ESHS response of HA solutions. The initial rise in the intensity with HA concentration is due to the increasing number density of the number of HA chains and their extended hydration shells. At the same time, increasing the HA concentration (in micrograms per milliliter). It can be seen that the radius of gyration \( r_g \) and the free radius per HA chain \( R_{free} \). (B) Temperature dependence of total fs-ESHS intensity and (C) maximum scattering angle at HA concentrations (molar weight, 1350 kDa) of 0.5 \( \mu \)g/ml (red circles) and 5 \( \mu \)g/ml (blue squares).
range. Because OSF theory is valid only for rigid polyelectrolytes such as DNA, i.e., when \( a \ll \sqrt{\lambda_d} L_0 \) (20), \( R_g \) is likely overestimated. Furthermore, because the contrast mechanisms vary between linear and nonlinear scattering, making a direct comparison of size parameters is not possible. Thus, while the size determined by nonlinear light scattering is of the same order of magnitude compared to what could be found from linear light scattering measurement if the experiment were possible, due to the number of uncertainties, it is very difficult to make any more precise conclusions.

**Extended hydration shell**

Last, we determine the extended hydration shell (\( R_h \)) around the polyelectrolyte. The extended hydration shell is determined by the water that interacts with the HA chain and hence does not relate directly to \( R \) or \( R' \). The maximum extended hydration shell that can be reached theoretically is determined by the available free volume per HA molecule. Figure 4A also shows the radius based on the available free volume (red line, \( R_{\text{free}} \)). When \( R_{\text{free}} > R' \), there is no interaction between the HA chains or soft shells [\( c_{\text{HA}} < 0.22 \text{ nM} \) (0.3 \( \mu \text{g/ml} \)]. When \( R_{\text{free}} < R' \), we expect that the extended hydration shell will be in the range \( R < R_h < R' \). At the concentration where the maximum intensity is reached, we expect that the extended hydration shell is maximum. To determine its magnitude, instead of using a model, we have performed fs-ESHS experiments at different temperatures, which allows us to vary the interaction strength of the HA chains and water and thus determine whether \( R_{\text{free}} = R_h \).

Increasing the temperature increases the relative importance of random thermal motion compared to ion-water (charge-dipole) interactions. It also changes the screening length. With increasing temperature, the orientational order of water will reduce, and we expect that this will lead to a decrease in the integrated fs-ESHS intensity \( S_{\text{int}} \). This intensity decrease arises from both a change in orientational order and a potential change in \( R \) due to the altered dipole-charge interactions. If the effective size \( R \) decreases with temperature as well, we expect to see an increase in the scattering angle of maximum intensity \( \theta_{\text{max}} \). However, if \( R \) is limited by the available free volume per HA chain (\( R_{\text{free}} \)), we do not expect to see an additional change in \( \theta_{\text{max}} \). In this case, the size of the hydration shell is equal to the radius of free volume per HA chain shown in Fig. 4A, i.e., \( R_{\text{free}} = R_h \).

Figure 4B shows the temperature dependence of the total fs-ESHS intensity measured between 0° and 100°C at two HA concentrations: 0.371 (0.5 \( \mu \text{g/ml} \)) and 3.71 nM (5 \( \mu \text{g/ml} \)), indicated by the dashed lines in Fig. 4A. As expected, the intensity decreases monotonically with temperature due to decreased orientational order. Note that the increase of temperature from 0° to 100°C also increases the Debye length by 20 nm at \( c_{\text{HA}} = 0.371 \text{ nM} \) (0.5 \( \mu \text{g/ml} \)) and by 15 nm at \( c_{\text{HA}} = 3.71 \text{ nM} \) (5 \( \mu \text{g/ml} \)), but this effect is not significant compared to the decrease in the orientational order from thermal motion. The maximum scattering angle \( \theta_{\text{max}} \) as a function of temperature for the two HA concentrations is plotted in Fig. 4C. At \( c_{\text{HA}} = 0.371 \text{ nM} \) (0.5 \( \mu \text{g/ml} \)), we observe a rise in \( \theta_{\text{max}} \) with temperature that reflects the shrinking of the extended hydration shell, and thus, the extended hydration shell is smaller than the free radius. However, at \( c_{\text{HA}} = 3.71 \text{ nM} \) (5 \( \mu \text{g/ml} \)), \( \theta_{\text{max}} \) is independent of temperature, indicating that the size of the hydration shell is constant. We therefore conclude that the theoretical extent of the hydration shells is at least equal to the free radius of HA at 3.71 nM, i.e., \( R_h \geq R_{\text{free}} \), amounting to an extended hydration shell with a radius of 27 ± 4 nm.

A shell with a radius of 475 nm holds \( ~1.3 \times 10^{10} \) water molecules and one polymer chain, whose volume will displace only \( ~20,000 \) water molecules. The measured value of \( R_b \), therefore reports on spatiotemporally averaged values. Compared to the value found for \( R \) (458 nm), which represents the radial position where the electrostatic potential changes, there is thus an additional shell of oriented water that is associated with the HA chain that has a thickness of 17 nm (37% of the Debye length at this concentration) or ~60 water molecules in diameter. This number of hydrating water molecules that surround the charged HA polymer is similar to the number of ~77 water molecules that were found around electrolytes in an earlier fs-ESHS study (27).

In summary, we have probed the extended hydration of HA, a biological anionic polyelectrolyte, using fs-ESHS. The source of the measured coherent second-harmonic radiation is assigned to water molecules that interact with the electrostatic field of spatially correlated charges on the polyelectrolyte chain. The fs-ESHS response is compared to dilute electrolyte solutions, which also display an increase in second-harmonic intensity at very low ionic strengths (~10^{-5} M). Unlike for simple electrolyte solutions, the fs-ESHS intensity versus HA concentration curve displays a maximum at an intermediate concentration (~13.3 \( \mu \text{g/ml} \), 11.6 nM) that is 6× larger than the intensity scattered by simple electrolytes. This difference is explained by the larger magnitude of the electrostatic field generated by HA compared to randomly distributed ions. At higher concentrations, the intensities scattered by HA and electrolyte solutions converge to the same plateau. This is explained by an enhancement of Debye screening with increasing HA concentration that results in a shrinkage of the hydration shells. A theoretical model that treats HA as a flexible charged particle surrounded by a spherical shell of polarized water molecules successfully describes the changes in the fs-ESHS intensity and angle as a function of HA concentration (ionic strength).

Using this model, we propose a method to determine the size of polyelectrolyte hydrates in dilute solutions. Temperature-dependent fs-ESHS experiments were used to estimate the size of the extended hydration shell, which amounts to a few hundred nanometers for a concentration of 13.3 \( \mu \text{g/ml} \) (11.6 nM). Similar to simple salt solutions, a clear difference between light and heavy water is observed. In heavy water, the fs-ESHS intensity maximum is 4.3× lower and occurs at 2.4× higher HA concentration (ionic strength). This difference cannot be explained by the proposed model and likely stems from the same nuclear quantum effect observed in simple electrolyte solutions (27). This isotope effect indicates that the interactions leading to the observed behavior not only are charge-water interactions but also contain interactions of the electrostatic field with the collective hydrogen bond network.

**MATERIALS AND METHODS**

**Chemicals**

HA (sodium hyaluronate, structure shown in Fig. 1A) produced by microbial fermentation of *Streptococcus pyogenes* was purchased...
from R&D Systems (molecular weight, 1350 kDa) and Contipro a.s. (molecular weight, 1145 kDa) and used as received. The salts NaCl (Sigma-Aldrich; >99.999%) and MgCl2 (Chempura; >99.995%) were filtered through Millipore Millex-VV 0.1-μm polyvinylidene difluoride membrane filters. We used ultrapure water (>18.2 megohm-cm) dispensed from a Milli-Q UF-Plus instrument (Millipore Inc.). Ultrapure heavy water (D2O, 99.8% D) was purchased from Armar Chemicals. Stock solutions of HA were prepared by reconstituting a weighted amount of HA powder in water. Individual samples were prepared by diluting a stock solution of HA in new Eppendorf tubes. The tubes were cleaned before use by rinsing with water.

**Determination of the ionic strength at infinite dilution**

The ionic strength at infinite dilution, I0, was estimated from conductivity measurements with an HI 5522 pH/ISE/EC bench meter and HI 76312 conductivity electrode (Hanna Instruments) in combination with tabulated ionic molar conductivities (73) using the formula

\[
I_0 = \frac{\kappa}{\sum_i \lambda_i \nu_i}
\]

where \(\kappa\) is the specific conductance, \(\lambda_i\) is the equivalent ionic conductivities of the cations and anions, and \(\nu_i\) refers to the number of moles of cations and anions. To estimate \(I_0\), we measured the ionic strength of pure water that was contained in the SHS cuvettes. On the basis of these conductivity measurements, we determined \(I_0\) = (5.1 ± 2.3) \times 10^{-6} M, assuming monovalent ions with \(\lambda_i\) = 60 \times 10^{-4} S m^2 mol^{-1} (73).

**Angle-resolved second-harmonic scattering**

The light source for fs-ESHS measurements was a Yb:KGW laser (Pharos SP, Light Conversion) producing 190-fs pulses centered at 1028 nm with a 200-kHz repetition rate. The polarization of the incident pulses was controlled via a Glan-Taylor polarizer (GT10-B, Thorlabs) and a zero-order half-wave plate (WPH05M-1030, Thorlabs). The incident laser pulses were filtered via a long-pass filter with a cutoff wavelength at 750 nm (FEL0750, Thorlabs). The average incident power at the sample was 60 mW (0.3 μJ per pulse). The laser beam was focused into a disposable cylindrical glass cuvette with an inner diameter of 4.2 mm (LS Instruments). The beam waist in the sample was 35 μm at the focus with a Rayleigh length of 0.94 mm. SHS light at 514 nm was collected with a plano-convex lens (f = 50 mm) and then filtered through a 10- or 50-nm-wide bandpass filter centered at 515 nm (ET515/10; ET515/50, Chroma). A Glan-Taylor polarizer (GT10-A, Thorlabs) was used for the polarization analysis of the scattered light. The polarization of the incident and outgoing light was either parallel (P) or perpendicular (S) with respect to the horizontal scattering plane. The SH light was focused using a plano-convex lens (f = 25 mm) into a gated photomultiplier tube (H7421-40, Hamamatsu). A more detailed description of the setup can be found in (74).

The experimental conditions used for fixed-angle measurements in Fig. 1 and figs. S2 and S4 were similar to those of (27, 48, 52). The detection angle was set to 90° with an acceptance angle of 11.4°. Each data point is an average of three to five measurements. Each measurement is an average of 50 exposures lasting 1 s each, i.e., using 50 \times 2 \times 10^4 pulses in total. The gate width was 10 ns. The fs-ESHS intensity at 90° was normalized by dividing the measured intensity of the sample solution by the intensity of the neat solvent (H2O or D2O) in the same polarization combination.

Angle-resolved fs-ESHS measurements shown in Figs. 2 and 4 and figs. S1 and S3 were performed by moving the detector arm in 5° steps between −90° and +90° with an opening angle of 3.4°. The normalized intensity at the angle \(\theta\), \(S(\theta)\), was normalized to the intensity of the neat water (H2O or D2O) using the formula

\[
S(\theta) = \frac{I(\theta)_{\text{ppp, sample}} - I(\theta)_{\text{ppp, water}}}{I(\theta)_{\text{SSS, water}}}
\]

where PPPP and SSS refer to the polarization combination of the incident and outgoing light [the first letter refers to the polarization state of the SH beam, and the second and third letters refer to that of the fundamental beam; P (S) is parallel (perpendicular) to the scattering plane]. The integrated fs-ESHS intensity was calculated by summing the normalized fs-ESHS intensity over all angles between −90° and 90° (except for 0°) using the formula

\[
S_{\text{int}} = \sum_{\theta \neq 0} S(\theta)
\]

**SUPPLEMENTARY MATERIALS**

Supplementary material for this article is available at http://advances.sciencemag.org/cgi/content/full/7/10/eabf2558/DC1

View/request a protocol for this paper from Bio-protocol.

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H.I.O., and S.R. interpreted the data and wrote the manuscript. S.R. conceived and supervised the work. The supporting material contains the following: fig. S1, power dependence of the fs-ESHS response; fig. S2, spectral recording of the fs-ESHS response; fig. S3, second-harmonic scattering patterns for HA (molecular weight, 1349 kDa) at various concentrations; fig. S4, normalized second-harmonic scattering intensity measured at a fixed angle of 90° with SSS polarization combination; fig. S5, fs-ESHS response from HA solutions at constant HA concentration as a function of excess ionic strength from added salt; fig. S6: $S_{ss}$ and $\theta_{max}$ for other polyelectrolytes as well as a neutral polymer; table S1, experimental size studies in the literature of HA in aqueous solution. Competing interests: The authors declare that they have no competing interests. Data and materials availability: All data needed to evaluate the conclusions in the paper are present in the paper and/or the Supplementary Materials. Additional data related to this paper may be requested from the authors.

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