

Dynamic Light Scattering Microrheology Reveals Multiscale Viscoelasticity of Polymer Gels and Precious Biological Materials

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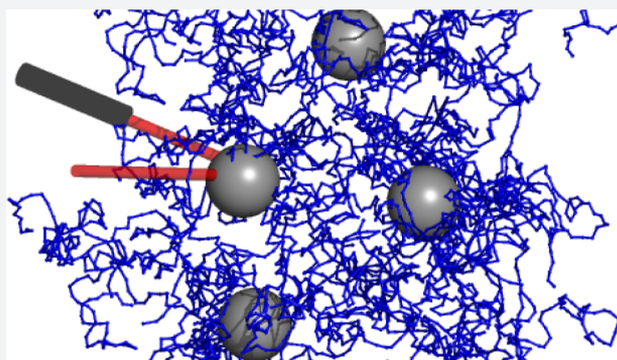
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Supporting Information

ABSTRACT: The development of experimental techniques capable of probing the viscoelasticity of soft materials over a broad range of time scales is essential to uncovering the physics that governs their behavior. In this work, we develop a microrheology technique that requires only 12 μL of sample and is capable of resolving dynamic behavior ranging in time scales from 10^{-6} to 10 s. Our approach, based on dynamic light scattering in the single-scattering limit, enables the study of polymer gels and other soft materials over a vastly larger hierarchy of time scales than macrorheology measurements. Our technique captures the viscoelastic modulus of polymer hydrogels with a broad range of stiffnesses from 10 to 10^4 Pa. We harness these capabilities to capture hierarchical molecular relaxations in DNA and to study the rheology of precious biological materials that are impractical for macrorheology measurements, including decellularized extracellular matrices and intestinal mucus. The use of a commercially available benchtop setup that is already available to a variety of soft matter researchers renders microrheology measurements accessible to a broader range of users than existing techniques, with the potential to reveal the physics that underlies complex polymer hydrogels and biological materials.



INTRODUCTION

Soft materials exhibit a rich range of rheological behaviors that play a central role in determining their processing behavior and practical functions. Examples include the engineered materials for biomedical applications,¹ self-healing materials for synthetic skin and flexible electronics,² self-assembling copolymers for nanopatterning,³ and the active cytoskeleton of living cells.⁴ The rheology of soft materials is governed by physical processes that occur over a vast range of time scales,⁵ which presents a formidable challenge for unlocking the molecular underpinnings of viscoelastic behavior. Thus, the development of rheological measurement techniques that interrogate the viscoelasticity of soft materials across disparate time scales offers the opportunity to gain deep insights into the molecular physics of soft matter with broad applications across scientific, engineering, and medical disciplines. Despite significant progress that has been forged in developing microrheology techniques that surpass the capabilities of conventional rheology and are amenable to the small sample quantities often encountered for biological materials,^{6–8} the requirements for specialized expertise, time-consuming data acquisition and

analysis, or a limited range of detectable material properties pose substantial barriers to establishing these techniques as accessible characterization tools across a broader community of materials chemistry and biomaterials researchers.

Conventional rheological techniques (macrorheology), which measure the material response to macroscopic external perturbations, typically provide access to a very limited range of time scales. In particular, oscillatory rheology provides access to frequencies ω of only up to about 10^2 s^{-1} . Although broader ranges of time scales can be accessed by combining oscillatory shear rheology with squeeze-flow and torsional resonance techniques, this requires the combined use of three separate measurements to access the frequency range of interest.^{9,10} Furthermore, such macrorheology experiments typically require large volumes of material, which prohibits measurements on certain precious biological samples, where only microliter-scale volumes can be obtained and procurement is often time-consuming and costly.

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To surmount these challenges, a variety of microrheology techniques have emerged that probe the viscoelasticity of soft matter at time scales and sample volumes beyond the reach of conventional macrorheology.^{6,7,11,12} Microrheology techniques typically employ micrometer-scale probe particles that sense their local viscoelastic environment in response to thermal (passive) or external (active) forces. Among these techniques, video particle tracking (VPT)^{7,13,14} and quadrant detection^{8,12,15–17} microrheology have enjoyed extensive implementation for probing spatially localized viscoelasticity in heterogeneous samples and living cells and require very low sample volumes (<1 μL). Spatially localized information from these techniques is achieved either by explicitly tracking particle trajectories by optical microscopy (VPT) or by focusing a laser onto individual probe particles and monitoring laser deflections onto a quadrant photodiode detector (quadrant detection). For bulk-averaged viscoelasticity across broad time scales, dynamic light scattering in the multiple-scattering limit, i.e., diffusing wave spectroscopy (DWS), has also been widely leveraged.^{11,15,18} However, each of these techniques suffers from distinct requirements that limit their widespread adoption by users beyond microrheology specialists, such as applicability to only soft materials ($G < 100$ Pa for VPT¹⁹), technically challenging implementation (quadrant detection and VPT), low statistical power (quadrant detection), or large sample volume requirements (at least 150 μL for DWS^{6,20–27}). Thus, there exists a critical need for a technique that is readily available to researchers with a broad range of expertise and that measures precious soft and biological materials across a breadth of viscoelastic properties and time scales.

Here, we develop a technique that is broadly accessible to users with a range of expertise and that interrogates viscoelasticity of small-volume samples over a vast spectrum of material properties and time scales. Our experimental methodology is based on dynamic light scattering (DLS) in the single-scattering limit and hence requires only dilute probe concentrations (<0.5%). This DLS microrheology (DLS μR) technique extracts the frequency-dependent shear moduli of materials with a broad range of stiffnesses up to 10^4 Pa and over a vastly broader range of time scales (10^{-1} to 10^6 s⁻¹) than oscillatory macrorheology (< 10^2 s⁻¹). This is achieved using a commercial benchtop instrument that is already available to a variety of materials chemists, involves minimal user intervention, and requires sample volumes as low as 12 μL . We leverage this technique to measure the hierarchy of molecular relaxations that occur in DNA solutions and to capture the viscoelastic behavior of precious biological materials, including extracellular matrices and mucus, where conventional macrorheology is impractical (Figure 1). This technique is accessible to a wide range of users and opens opportunities for studying a range of complex biological and soft materials.

The ranges of frequency and viscoelasticity that we explore in this work have typically been viewed to be inaccessible to DLS in the single-scattering limit.^{18,28–31} However, as we demonstrate, this perceived limitation reflects misconceptions related to the statistics of photon correlation in single-scattering detection. We harness the highest spatial resolution accessible to single-scattering DLS by operating in noninvasive backscatter detection mode, and prove that commercial photon correlation instruments are capable of capturing probe fluctuations at nanometer-scale resolution. This reveals access to time scales and material stiffness previously viewed to be

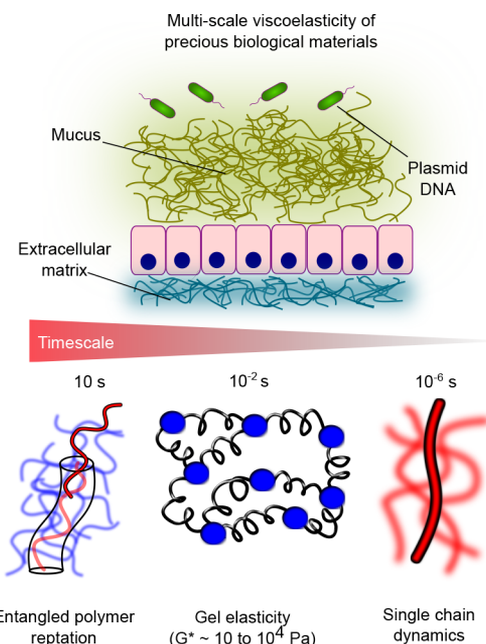


Figure 1. DLS μR captures the multiscale viscoelasticity of soft matter, including polymeric gels and precious biological materials, such as DNA solutions, intestinal mucus, and extracellular matrices. DLS μR reveals signatures of molecular relaxations that occur over 7 decades in time, ranging from 10 s down to 10^{-6} s, which spans the time scales associated with reptation of polymers in entangled networks, entropic elasticity of cross-linked or entangled gels, and internal relaxations of individual polymer chains.

accessible by microrheology only through DWS and quadrant detection.

RESULTS

DLS μR Offers an Accessible Technique for Rheological Measurements of Soft Matter over a Wide Spectrum of Time Scales and Material Properties. Figure 2 illustrates our workflow for extracting the frequency-dependent shear modulus $G^*(\omega)$ of soft materials using a commercially available DLS instrument. In contrast to DWS, by operating in the single-scattering limit, our methodology involves dilute

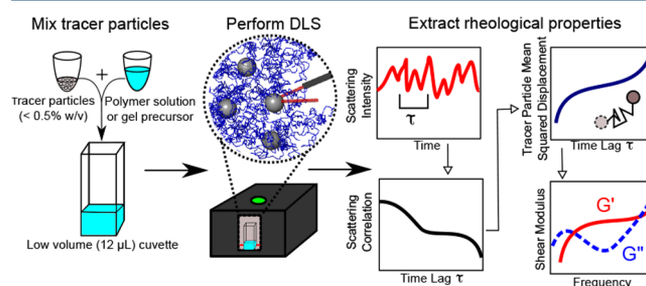


Figure 2. DLS μR workflow. The polymer solution or gel precursor is mixed with a dilute concentration of tracer particles (<0.5% w/v). DLS is performed in a backscattering configuration using a commercial benchtop instrument. Brownian motion of the tracer particles produces fluctuations in scattering intensity that give rise to a characteristic scattering intensity autocorrelation. The autocorrelation is analyzed by our custom software to extract the mean-squared displacement of particles, which is used to determine the frequency-dependent linear viscoelastic shear modulus $G^*(\omega)$.

concentrations of probe particles (<0.5% w/v) that serve as light scatterers.

The commercial benchtop instrument we leverage here illuminates the sample with a 50 μm diameter laser beam, and scattered light is focused onto a photodiode detector using a translating lens to collect photons from a specified scattering volume within the sample of about 1.4 nL. In practice, we use a total sample volume of 12 μL . This strikingly contrasts with the volume requirements of DWS, where at least 150 μL of sample is typically required.^{6,20–27} We perform DLS in backscatter detection mode, which offers the smallest length scale of resolution accessible to the wavelength of the laser, thereby capturing time scales that are inaccessible to oscillatory shear rheology and VPT. In contrast to DWS, the low probe concentrations typically required for performing single-scattering DLS can present a challenge for separating the scattering by probe particles from background scattering by the material.³² Another key advantage of the backscattering optics utilized here is that much higher probe concentrations can be used without exiting the single-scattering regime due to the shorter photon path length in the backscattering configuration and the ability to tune the photon path length using the translating lens system.³³ Leveraging these capabilities enables us to use probe concentrations (0.1% to 0.25%) that are 1–2 orders of magnitude greater than those typically implemented for single-scattering DLS,^{18,31,32} thereby ensuring dominance of scattering from probe particles.

Thermal fluctuations of the tracer particles give rise to scattering intensity fluctuations, which are described by a time lag τ dependent scattering intensity autocorrelation $g^{(2)}(\tau)$. Scattering photon autocorrelations are collected using a digital correlator with delay times ranging from 0.5 μs to 70 s, and the experimentally accessible frequency range within this window is dictated by the time scales over which particle displacements are detectable. The intensity autocorrelation encodes the average mean-squared displacement of tracer particles within the scattering volume $\langle \Delta r^2(\tau) \rangle$ over the time lag τ . The mean-squared displacement of particles with radius a in turn can be used to extract the frequency-dependent shear modulus according to the generalized Stokes–Einstein relation, $G^*(\omega) = k_B T / [\pi a i \omega \langle \Delta r^2(\omega) \rangle]$. Our custom software package analyzes raw scattering intensity autocorrelations to extract $G^*(\omega)$ of the material.

To investigate the capability of our technique to capture the viscoelastic behavior of polymer gels, we study chemically cross-linked polyacrylamide networks over a broad range of stiffnesses (shear moduli G^* from 10 to 10^4 Pa) and compare our DLS μR measurements to those obtained from conventional oscillatory macrorheology. Over this full range of stiffnesses, our DLS μR measurements exhibit excellent agreement with macrorheology in the frequency range in which the two techniques overlap (Figure 3). This viscoelasticity spectrum accessible to DLS μR spans the stiffnesses of a diverse range of biological tissues, ranging from mucus to skeletal muscle.³⁴ This range of biologically relevant stiffnesses is inaccessible to VPT, which would be limited to materials with $|G^*| < 10^2$ Pa using the 100 nm diameter probe particles used in our experiments.¹⁹ We note that although generally good agreement is found between our DLS μR measurements and macrorheology, the modulus in our DLS μR experiments underestimates the modulus by about a factor of 2 for the stiffer (5% and 10%) gels. This may be due to inhomogeneities formed during the free-radical polymerization, which are likely to be more

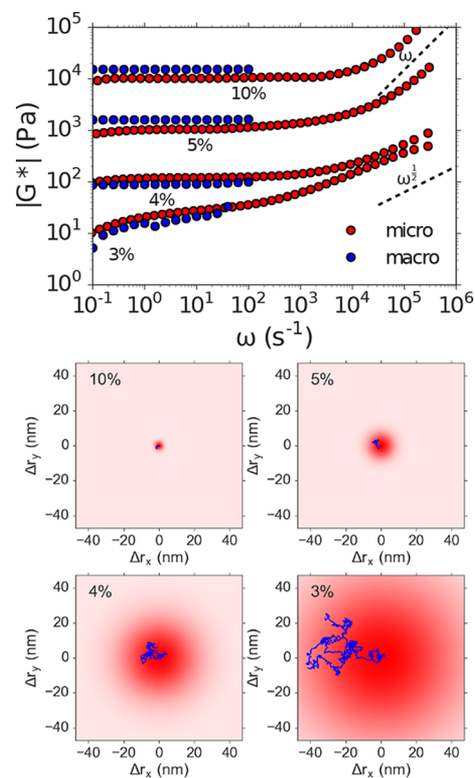


Figure 3. DLS μR recapitulates macrorheology in cross-linked polyacrylamide gels with shear moduli G^* spanning 10^1 to 10^4 Pa. Top: Comparison of the frequency ω dependence of the magnitude of the shear modulus $|G^*|$ obtained by DLS μR and macrorheology of polyacrylamide gels with varying polyacrylamide concentrations (% w/v). The dashed lines represent the high-frequency scaling of a Rouse polymer $G^* \sim \omega^{1/2}$ and space-filling branched polymer fractals with nondraining hydrodynamics $G^* \sim \omega$. Bottom: Illustration of typical particle trajectories occurring over a time interval corresponding to the angular frequency $\omega = 10^1$ s for a 100 nm diameter tracer particle embedded in gels with the indicated polyacrylamide composition.

prevalent in the more rapidly forming gels.³⁵ Such inhomogeneities may force the probe particles into more compliant regions of the network. A similar effect of inhomogeneities in gel microrheology has been previously described,⁷ and particle tracking microrheology experiments have demonstrated the existence of mechanical heterogeneity in polyacrylamide gels.³¹

Moreover, DLS μR captures a vastly broader range of time scales than conventional oscillatory macrorheology and VPT, ranging from frequencies ω of 10^{-1} to 10^6 s^{-1} . Although similar frequency ranges have been captured in macrorheology measurements that combine oscillatory rheology, squeeze-flow, and torsional resonance, this is achieved only with the combined use of 3 separate measurements, in total requiring hundreds of microliters of material.^{9,10} Our technique enables us to probe a hierarchy of viscoelastic processes in the material in a single measurement with only 12 μL of sample. Our measurements demonstrate that, at intermediate to long time scales, the gels behave as elastic solids in which G^* is approximately constant with respect to frequency. At short time scales, the tracer particles probe faster relaxation modes in the gel due to effective elastic chains that give rise to a power-law scaling with $G^* \sim \omega^\alpha$. Consistent with the decreasing molecular weight of effective elastic chains with increasing cross-linking density, we find that the time scale at which these relaxation modes emerge decreases by 2 orders of magnitude as the gel

stiffness increases across the range spanned in our measurements (Supporting Information and Supporting Figure 1).

For the most compliant gels, $\alpha \approx 1/2$, consistent with freely draining Rouse dynamics of flexible partial chains between cross-links.³⁶ This suggests that, at low polymer concentrations, the gel network is formed from loosely cross-linked high molecular weight polymers in which each effective elastic chain experiences overlap with surrounding chains that screen hydrodynamic interactions. For the more densely cross-linked gels, the high-frequency scaling behavior approaches $\alpha \approx 1$, which coincides with the critical scaling predicted by percolation theory at the gel point for nondraining percolating clusters³⁷ as well as the expected scaling behavior for nondraining Zimm relaxation of self-similar branched chains with a fractal dimension $D_f = 3$.³⁸ This relaxation can be interpreted in terms of a network that consists of effective elastic chains that possess an internal fractal branched structure that is space-filling at all length scales below the gel correlation length and retains high-frequency signatures of the fractal structure and dynamics present at the percolation point (see Supporting Information for derivation). This is consistent with time–temperature superposition experiments in which fully formed gels often exhibit scaling of G'' that mirrors the fractal dynamics at the gel point.³⁹

The transition in the high-frequency relaxation from Rouse to critically overlapped Zimm dynamics suggests that the greater cross-linking efficiency in the higher concentration gels produces more tightly cross-linked networks consisting of branched effective elastic chains that do not overlap with surrounding chains, and therefore do not experience screening of hydrodynamic interactions. This is consistent with time-cure superposition experiments just above the gel point, which show larger critical relaxation scaling exponents with greater cross-linking efficiency (compare ref 40 and ref 41) or lower molecular weight polymer precursors.⁴² Similarly, atomic force microscopy microrheology experiments on polyacrylamide gels at frequencies up to $\omega \approx 10^3$ rad/s have demonstrated a high-frequency scaling exponent that increases with polyacrylamide content, approaching $\alpha \approx 1$ for 10 kPa gels.⁴³ Our technique thus offers a substantial advantage over oscillatory shear rheology in terms of elucidating the hierarchy of physical processes that govern the behavior of polymer gels.

Unlike VPT and quadrant detection microrheology, which involve explicit tracking of particle trajectories, DLS μ R yields measurements of tracer particle fluctuations that are directly averaged over the 1.4 nL scattering volume. Hence, VPT and quadrant detection extract spatially heterogeneous viscoelasticity that is concealed by averaging inherent to DLS μ R. However, our technique offers a more streamlined and accessible work flow for diverse users than VPT and quadrant detection microrheology. In contrast to VPT, in which each time point requires a full microscopy image of tracer particle positions to be acquired, stored, and processed, DLS μ R measurements directly provide the statistically averaged particle fluctuations in the form of the scattering intensity autocorrelation function, thereby reducing the data footprint associated with each measurement by several orders of magnitude. Unlike VPT, quadrant detection can access the range of viscoelasticity and time scales that are encompassed by DLS μ R, but quadrant detection requires tracking individual trajectories at a time by optically trapping single probe particles. This comes at the expense of vastly decreased statistical power, even compared to VPT, in which approximately 100 particles are typically present

in the field of view. Moreover, quadrant detection microrheology requires expertise with optical traps that is not commonly found in many soft materials laboratories. Together, these limitations render quadrant detection a far less accessible and more technically challenging approach. By comparison, DLS μ R is readily implemented on a benchtop commercial instrument that is already available to many materials chemistry and biomaterials researchers and can rapidly capture viscoelasticity with high statistical power while requiring minimal user intervention.

Historically, DLS μ R in the single-scattering limit has found limited use in comparison to VPT, quadrant detection, and DWS, and it is conventionally considered to be inadequate for viscoelastic properties in the stiffness and time-scale ranges probed in our measurements.^{18,28–31} By comparison to quadrant detection and DWS, previous studies in single-scattering detection have typically explored a much more limited range of stiffnesses ($G^* < 100$ Pa) and frequencies $\omega < 10^4$ s⁻¹.^{18,31,32,44–46} The range of stiffnesses and time scales accessible is fundamentally limited by the length scale of tracer particle fluctuations to which the light-scattering configuration is sensitive (visualized in Figure 3). This can be understood on the basis of the generalized Stokes–Einstein relation, wherein the mean-squared displacements of tracer particles in an elastic gel are inversely proportional to the shear modulus.¹¹ Thus, the sensitivity is dictated by the smallest length scale of probe fluctuations that produce detectable decreases in the scattering intensity autocorrelation $g^{(2)}(\tau)$.

We find that the backscattering optics utilized here are capable of resolving decreases in the scattering intensity autocorrelation that correspond to mean-squared displacements of about 1 nm², which are 2 orders of magnitude smaller than the mean-squared displacements detectable by VPT,¹⁹ as well as those previously accessed using single-scattering DLS for microrheology of soft polyacrylamide gels.³¹ In accordance with the generalized Stokes–Einstein relation, the minimum detectable displacement, together with the probe particle size, dictates the maximum measurable stiffness. The 1 nm resolution achieved here renders measurements on gels as stiff as 10⁴ Pa possible using 100 nm beads, and the maximum accessible stiffness scales inversely with the size of the probe that is required to satisfy the continuum assumption of the Stokes–Einstein relation. In our experiments, the 100 nm beads are much larger than the mesh size of the polyacrylamide network, which will be about 10 nm or less in the range of concentrations we probe,⁴⁷ and are therefore sufficiently large to ensure that the continuum limit is reached.

It is noteworthy that the length scale of fluctuations to which we are sensitive is much smaller than the length scale that would be estimated based on the scattering wave vector q . For the backscattering angle θ (173°), the wavelength λ of the laser (633 nm), and a refractive index n equal to that of water, the scattering wave vector in our experiments $q = 4\pi n \sin(\theta/2)/\lambda$ corresponds to a length scale $q^{-1} = 38$ nm.⁴⁸ For this reason, DLS μ R in single-scattering detection has often been viewed as unsuitable for measurements in the stiffness and frequency ranges that require the spatial resolution captured in our experiments.^{18,28} In addition to improved spatial resolution through the use of backscattering optics compared to some previous studies using forward or orthogonal scattering,^{18,31} higher spatial resolution is also achieved by accurate estimation of the correlation function zero-time intercept using the methods described in the Supporting Information. Accurate

estimation of the zero-time correlation intercept has been previously reported to represent a limitation for DLS microrheology.¹⁸ Our results emphasize the importance of considering the sensitivity of photon correlation detection and the reliability of correlation intercept estimation in evaluating the limits of the DLS μ R technique.

Polymer gels exhibit broken ergodicity due to frozen-in density fluctuations formed during the gelation process, which must be accounted for in interpretation of DLS autocorrelation data.⁴⁹ In principle, broken ergodicity can be corrected for by obtaining ensemble-averaged scattering intensity autocorrelation functions by collecting the time-averaged autocorrelation across an ensemble of spatial positions in the sample, which is time-consuming and tedious. The broken-ergodicity correction we implement here (described in the [Supporting Information](#)) enables extraction of the contribution to the scattering intensity due to dynamic fluctuations within the scattering volume of interest, while obviating the need for full ensemble averaging of the correlation function. However, we note that the acquisition times we use here to collect high-quality photon statistics are not amenable to rapidly evolving materials, such as during the gelation process.

DLS μ R Elucidates Hierarchical Molecular Relaxations in DNA. The broad spectrum of time scales probed by DLS offers the opportunity to gain insights into the physics of macromolecules that exhibit distinct relaxation behaviors at different time scales. Semiflexible polymers constitute an important class of macromolecules that exhibit distinct physics at different length scales and time scales. The physical behavior of such polymers is governed by bending elasticity on the scale of their persistence length l_p , but they can exhibit flexible Gaussian chain behavior at lengths much longer than their persistence length. DNA represents an excellent model system for exploring this relaxation hierarchy, since its persistence length ($l_p \approx 50$ nm)⁵⁰ is sufficiently short to produce flexible Gaussian chain statistics at experimentally accessible molecular weights, but provides sufficient bending stiffness to yield semiflexible polymer behavior at experimentally accessible length scales and time scales.

DLS μ R measurements were performed on semidilute, unentangled solutions of linear DNA (1.0 mg/mL of 5.8 kilobase chains, i.e., contour length $L \approx 1.9$ μ m). Our results elucidate a hierarchy of regimes of physical behavior (Figure 4, designated as A, B, and C). We relate the transition between these regimes to two theoretically predicted time scales ($\tau \sim \omega^{-1}$): the bending relaxation time of semiflexible subsections of the DNA chain τ_{bend} and the Rouse time associated with total relaxation of all internal polymer elastic modes τ_R . These time scales approximately define three regimes that correspond at the shortest times ($\tau < \tau_{\text{bend}} < \tau_R$) to bending fluctuations in the semiflexible DNA double helix, at intermediate times ($\tau_{\text{bend}} < \tau < \tau_R$) to relaxations of internal segments of the chain that behave like flexible segments, and at the longest times ($\tau_{\text{bend}} < \tau_R < \tau$) to total chain diffusion. Our DLS μ R measurements agree quantitatively with theoretical predictions (dotted and solid curves in Figure 4) for a wormlike chain (WLC), which we obtain by combining the high-frequency scaling limit of a WLC with the full analytical solution of a Rouse polymer, as described in the [Supporting Information](#) and [Supporting Figure 5](#).

At the shortest times (largest ω , region C), we identify a rheological response which reflects thermal bending fluctuations in the DNA double helix due to its finite bending stiffness.

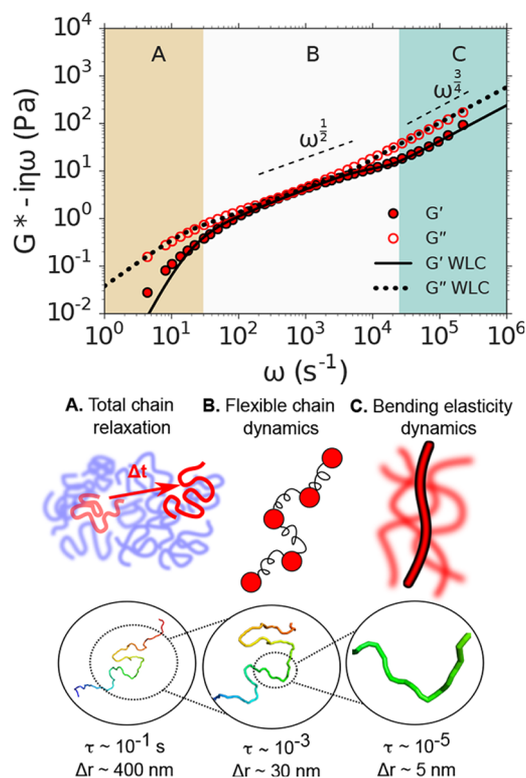


Figure 4. DLS μ R of DNA solutions reveals a hierarchy of molecular relaxations. Top: Shear modulus G^* as a function of angular frequency ω of semidilute DNA solutions. Regions A, B, and C represent approximate regimes in which the viscoelastic response is expected to probe the total chain relaxation, internal flexible chain relaxation, and bend relaxation, respectively. Dashed lines indicate the expected scaling laws for a Rouse polymer ($G^* \sim \omega^{1/2}$) and a freely draining semiflexible chain ($G^* \sim \omega^{3/4}$). The dotted and solid lines indicate the theoretical predictions for G' and G'' , respectively, of a semiflexible polymer modeled as a WLC. Middle: Illustrations of the physical behavior probed at the time-scale regimes annotated A, B, and C in the top plot. Bottom: Representative snapshot of a 5.8 kilobase DNA molecule modeled as a WLC obtained by Monte Carlo simulation. Magnifications represent the approximate contour length of the DNA molecule whose relaxations dominate the viscoelastic response that is probed at time scales τ (which reside within regimes A, B, and C) by tracer particles undergoing approximate displacements Δr .

In this regime, the scaling behavior in our DLS μ R measurements approaches the theoretical predictions for the high-frequency response of a WLC, in which the polymer contribution to the shear modulus is $G = \frac{2^{3/4}}{15} \rho (k_B T)^{1/4} l_p^{5/4} (i \omega \xi_{\perp})^{3/4}$,^{51,52} where ρ is the polymer contour length per unit volume, $k_B T$ is thermal energy, l_p is the persistence length, and ξ_{\perp} is the transverse friction coefficient per unit length (which we estimate to be 4×10^{-2} Pa \times s by fitting theoretical predictions for a WLC to our data using ξ_{\perp} as a fitting parameter). We theoretically estimate the time scale where these bending fluctuations will manifest to be $\tau_{\text{bend}} = \xi_{\perp} (2l_p)^4 / [(3\pi/2)^4 l_p k_B T] \approx 40$ μ s.⁵³ This theoretically predicted time scale (indicated by the boundary between regions B and C in Figure 4) corresponds approximately to the frequency $\omega = \tau_{\text{bend}}^{-1}$ near which the rheological behavior from DLS μ R transitions to the high-frequency scaling limit.

At intermediate time scales (region B), which are long in comparison to the relaxation time of bending modes but shorter than the time required for relaxation of all internal

polymer elastic modes, the shear modulus reflects the viscoelastic relaxation of flexible subchains that behave like entropic springs. For a Rouse-like polymer, in which hydrodynamic interactions can be neglected, this is theoretically predicted to produce a shear modulus in which the storage and loss moduli scale as $G' = G'' \sim \omega^{1/2}$.⁵ Our measurements show that the relaxation behavior of the DNA chains approaches this scaling behavior over a limited range of frequencies ($10^2 < \omega < 10^3$) before transitioning to total chain diffusion at longer time scales and bend fluctuation dynamics at shorter time scales.

At the longest time scales (smallest ω , region A), all internal modes of the polymer chain have fully relaxed, and the shear modulus captures the viscoelastic response due to translational diffusion of the polymer chains. For semidilute unentangled solutions, this response is anticipated at times longer than the Rouse relaxation time of the entire polymer contour length L : $\tau_R = 2l_p \xi_R L^2 / (3\pi^2 k_B T) \approx 3 \times 10^{-2}$ s.⁵ At frequencies $\omega < \tau_R^{-1}$, our measurements exhibit a transition to viscous dynamics, with $G'' > G'$, which is consistent with relaxation of the internal entropic elastic modes of the polymers and a transition toward total chain diffusion. This transition is in excellent agreement with theoretical predictions for a WLC that exhibits flexible chain behavior at these time scales, which we obtain using a Rouse monomer friction coefficient per unit length of $\xi_R = 1.0 \times 10^{-2}$ Pa \times s based on the fitting procedure described in [Supporting Information](#). The value of ξ_R we obtain is about 4 times smaller than the transverse friction coefficient ξ_{\perp} .

In order to evaluate the consistency of our fitted friction coefficients, we compare the value of ξ_R obtained from our DLS μ R data to existing diffusivity measurements and relate our observation that $\xi_{\perp} \approx 4\xi_R$ to polymer physics theory. In the semidilute regime, the effective Rouse friction coefficient ξ_R is dependent on the polymer concentration, which defines the length scale for hydrodynamic screening by overlapping polymers. At length scales larger than the hydrodynamic screening length, each polymer in the semidilute network can be viewed as consisting of coarse-grained polymer “blobs” that diffuse like effective Rouse monomers³⁶ and exhibit internal hydrodynamic interactions at smaller length and time scales. Measurements of the diffusivity D of fluorescently labeled 5.9 kb DNA have confirmed this concentration dependence and at a concentration of 1.0 mg/mL provide a Rouse friction coefficient $\xi_R = k_B T / (DL) = 0.6 \times 10^{-2}$ Pa \times s.^{54,55} This is somewhat lower than our value $\xi_R = 1.0 \times 10^{-2}$ Pa \times s, and further investigation is required to elucidate the origin of this discrepancy.

Below the hydrodynamic screening length, and within each effective Rouse monomer, DNA may be expected to behave according to slender body hydrodynamics.⁵⁶ In this regime, the hydrodynamics are described by two friction coefficients, ξ_{\perp} and ξ_{\parallel} , which represent the transverse and longitudinal friction, respectively, and $\xi_{\perp} = 2\xi_{\parallel}$. If the friction coefficient of each Rouse monomer corresponds to the total friction of each subsection that obeys slender body dynamics, then slender body theory implies that $\xi_R = 3(2/\xi_{\perp} + 1/\xi_{\parallel})^{-1} = 3\xi_{\perp}/4$. However, our fit to the DLS μ R data instead yields $\xi_R \approx \xi_{\perp}/4$. The origin of this disagreement is unclear, and may reflect additional many-body effects due to the semidilute solution not considered by slender body theory.

Importantly, our results are only slightly dependent on probe particle size for probe diameters of at least 500 nm (the probe size used in [Figure 4](#)), indicating that the probes sense the continuum viscoelasticity of the fluid ([Supporting Figure 2](#)).

This is consistent with previous microrheology studies of linear DNA solutions, which show that, at the concentration used in our work, and with DNA molecular weights where entanglements are weak or nonexistent, noncontinuum effects such as polymer depletion are not substantial, provided the probes are larger than the radius of gyration of the polymers.^{57,58} Unlike the polyacrylamide gels, 100 nm probes do not satisfy the continuum assumption of the Stokes–Einstein relation, since the radius of gyration of a 5.8 kilobase Gaussian DNA chain is about 180 nm, and the system resides close to the critical overlap concentration.

The transition to an $\omega^{3/4}$ scaling behavior at high frequencies has been previously demonstrated using quadrant detection and DWS microrheology of other semiflexible polymers, such as F-actin, wormlike micelles, and polysaccharides.^{8,59–62} However, despite the extensive history of DNA as a model polymer in the polymer physics community, previous microrheology studies on DNA solutions were conducted at much lower frequencies than the maximum frequency accessed here.^{6,63,64} Our measurements not only capture the expected high-frequency scaling behavior that has been confirmed in experiments on other semiflexible polymers but also exhibit quantitative agreement with polymer physics theory across the full range of molecular relaxations.

DLS μ R Accesses Multiscale Relaxation in Extracellular Matrices Where Time–Temperature Superposition Fails.

Time–temperature superposition is a widely implemented technique for broadening the frequency range that is accessed in a conventional macrorheology experiment. In brief, this technique involves collecting the frequency-dependent shear modulus at a range of temperatures and horizontally shifting the isotherms to construct a single “master curve” that encompasses a substantially broader range of frequencies than those accessed at any particular temperature.⁶⁵ However, this construction is valid only for materials in which the relaxation rates and physical processes that govern the viscoelastic behavior can be described by a single temperature dependence.

Biological systems provide a variety of instances in which the assumptions underlying time–temperature superposition fail. In fact, in general, time–temperature superposition does not substantially broaden the frequency range for aqueous polymer systems, since the available temperature variations do not lead to substantial frequency shift factors. In particular, extracellular matrices represent an important class of biological materials that exhibit nontrivial thermal responsiveness that violates time–temperature superposition. Furthermore, the rheological behavior of extracellular matrices exhibits a profound impact on directing cellular behavior.^{66,67}

To illustrate the utility of the DLS μ R for these materials, we conducted measurements on Matrigel across temperatures spanning its sol–gel transition. Matrigel consists of a complex mixture of extracellular matrix proteins and is widely used in mammalian tissue culture.⁶⁸ We find that DLS μ R provides excellent agreement with macrorheology at fixed temperature ([Supporting Figure 3](#)), but reveals a temperature-dependent viscoelastic spectrum that clearly violates time–temperature superposition ([Figure 5](#)). Below the sol–gel transition, Matrigel exhibits viscoelastic power-law behavior with $G'' > G'$ over the full range of accessible frequencies. Above the sol–gel transition, the shear modulus displays two distinct regimes consistent with a polymer hydrogel. At low to intermediate frequencies, a plateau modulus with a weak frequency dependence emerges with $G' > G''$, reflecting behavior

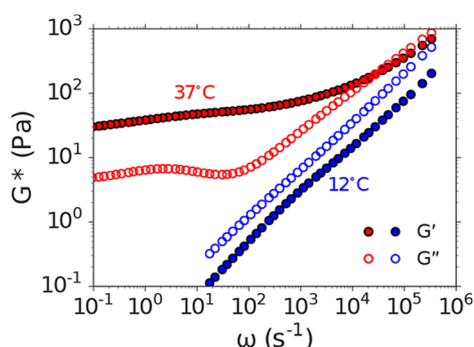


Figure 5. DLS μ R captures the viscoelastic response of Matrigel across a broad range of time scales where time–temperature superposition fails. Closed and open symbols represent frequency dependence ω of the storage and loss shear moduli (G' and G'' , respectively) at temperatures below (12° , blue) and above (37° , red) the sol–gel transition of Matrigel.

reminiscent of an elastic solid. At high frequencies ($\omega > 10^4$ s^{-1}), the shear modulus increases with frequency and approaches a power-law behavior that reflects coupling to the underlying internal relaxation modes. These results demonstrate that DLS μ R enables measurements of the broad frequency viscoelasticity of biological materials where conventional oscillatory rheology with time–temperature superposition is not an option.

DLS μ R Reveals Broad Frequency Entanglement Dynamics in Intestinal Mucus for Which Macrorheology Is Impractical.

Intestinal mucus lines the luminal surface of the intestinal epithelium and serves as an essential barrier whose functions include acting as a physical barrier against microbial invasion and providing nutritional support to the host microbiome.^{69–71} Disruptions to the mucosal layer are implicated in a variety of diseases, which suggests that the ability to quantify the physical properties of the mucosal layer may provide biophysical insights into the role of mucus in such pathologies. Intestinal mucus is available in only small quantities from small animal models and human patients, as only a small amount can be recovered from a single animal in a mouse model, and in human patients, mucus must be obtained from biopsies. Thus, intestinal mucus represents an excellent example of a biological material in which acquiring macro-rheology measurements is challenging, costly, and impractical.

Our DLS μ R measurements on intestinal mucus isolated from healthy mice (Figure 6) reveal a viscoelastic spectrum that spans 6 decades in time and exhibits signatures of a hydrogel formed by a network of physical entanglements among macromolecules. The frequency-dependent shear modulus reflects three distinct regimes of physical behavior that are predicted by the Doi and Edwards tube theory for entangled polymers. In this model, polymers within an entangled network are envisioned to be confined by the surrounding network within an effective tube that restricts their motion within the

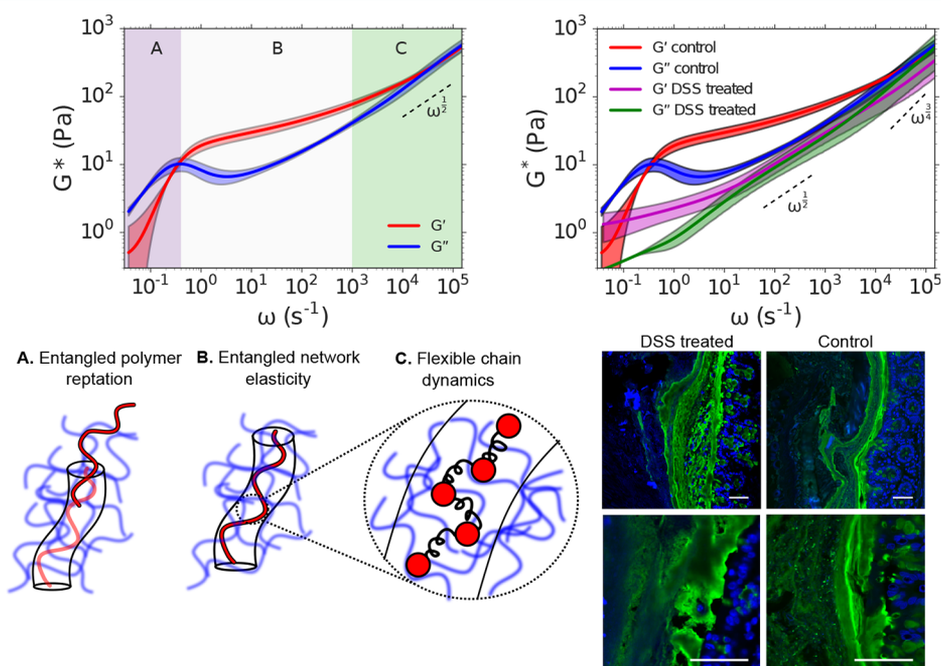


Figure 6. DLS μ R captures the entangled dynamics of intestinal mucus of healthy and colitic mice. Top left: Dependence of the shear modulus G^* on angular frequency ω of intestinal mucus isolated from healthy mice. The shear modulus exhibits three regimes, A, B, and C, which we identify as corresponding to reptation of polymers within an entangled network, elastic behavior due to entanglement constraints, and Rouse-like flexible chain dynamics at length scales below the entanglement confinement length, respectively. Solid curves represent the mean among 3 independent biological reproductions. Shading represents 90% confidence intervals of the mean generated by bootstrap resampling spectra from independent biological reproductions. The dashed line represents the high-frequency scaling behavior of a Rouse polymer $G^* \sim \omega^{1/2}$, which is provided to guide the eye. Bottom left: Illustration of the physical behavior probed in time-scale regimes annotated A, B, and C, as they are interpreted in terms of the Doi and Edwards tube model of entangled polymer dynamics. Top right: Comparison of the frequency-dependent shear modulus G^* of intestinal mucus isolated from healthy (control) mice and mice treated with dextran sulfate sodium (DSS) to induce colitis. The high frequency scaling behaviors of a Rouse polymer and a WLC ($G^* \sim \omega^{1/2}$ and $G^* \sim \omega^{3/4}$, respectively) are indicated with dashed lines for reference. Bottom right: Confocal imaging of DSS treated and control mouse colons recapitulates physical disruption of mucus. The intestinal sections were stained to highlight the mucus (UEA-1, green) and the DNA (DAPI, blue). In DSS treated mice the distal colon mucus loses much of the usual striated organization. Scale bars are 50 μ m.

tube diameter (see illustrations in Figure 6). At short times (large ω , region C), the viscoelastic spectrum is dominated by the internal relaxation of flexible subsections of the polymers that are smaller than the effective tube diameter and have yet to “discover” that they are confined by entanglements within the surrounding network. In this regime, $G^*(\omega)$ possesses a frequency scaling behavior that reflects single chain dynamics that are intermediate between a Rouse polymer and a WLC. At intermediate times (region B), polymer relaxations are governed by entanglement constraints with the surrounding polymer matrix, which produces an extended elastic plateau modulus in which $G' > G''$ and G' exhibits a weak frequency dependence. At sufficiently long times (small ω , region A), the polymers are able to escape their local entanglements by reptation, which enables the material to flow, yielding a viscous response with $G'' > G'$.

The rheological properties of intestinal mucus may play a pivotal role in gut barrier function as well as the growth of microbes embedded within the matrix.^{71,72} To demonstrate the capability of DLS μ R to capture changes in mucus rheology associated with loss of barrier function, we investigate intestinal mucus in a mouse colitis model. Incorporation of dextran sulfate sodium (DSS) into the drinking water of mice disrupts barrier function of intestinal mucus and induces inflammation of the underlying epithelium.⁷³ Figure 6 illustrates the change in mucus rheology between healthy and DSS treated (colitic) mice measured by DLS μ R. Our results exhibit a striking change in mucus rheology in the colitis model in which the mucus is substantially softened by DSS treatment, suggestive of a substantial reduction in the overall cross-linking density of the matrix (Figure 6). This perturbation to mucus viscoelasticity coincides with physical disruption of mucus morphology within the colon, as visualized by confocal microscopy (Figure 6). Furthermore, although we cannot obtain mucus from mice in sufficient quantities for macrorheology, to validate our DLS μ R measurements on mucus samples, we performed DLS μ R on reconstituted porcine gastric mucus and found quantitative agreement with macrorheology (Supporting Figure 4). These findings suggest that DLS μ R may serve as a valuable tool for investigating biophysical connections between intestinal mucus viscoelasticity and gut barrier status.

CONCLUSIONS

In this work, we present a microrheology technique based on DLS in the single-scattering limit that provides access to the frequency-dependent viscoelasticity of soft materials across up to 7 decades in time scale. This technique requires only 12 μ L of sample, which renders it amenable to precious materials where sample volume requirements for macrorheology and broad-frequency DWS are impractical. Moreover, we demonstrate that, contrary to common belief, DLS μ R in single-scattering detection mode is capable of probing the viscoelasticity of polymer hydrogels with a broad range of material properties, with shear moduli spanning 10 to 10⁴ Pa, which encompasses stiffnesses across a variety of biological tissues. Importantly, this technique is implemented with a commercial benchtop instrument and is therefore accessible to users with a broad range of expertise.

DLS μ R represents a valuable contribution to a suite of rheological tools that are available to researchers, and extends the domain of microrheology to a broader base of users. For applications where rapid characterization of ensemble-averaged viscoelasticity of precious materials is desired, DLS μ R provides

a powerful and accessible alternative to existing techniques. Although VPT and quadrant detection microrheology extract spatially heterogeneous viscoelasticity that is lost by ensemble averaging in DLS μ R, our technique overcomes crucial challenges in these established approaches that present barriers to their widespread implementation in the materials chemistry and biomaterials communities. By comparison to VPT, DLS μ R reveals viscoelasticity in materials 2 orders of magnitude stiffer and across a range of time scales that is 4 orders of magnitude greater than those accessible to VPT, while obviating the need for the large data footprint and cumbersome data analysis that is required of optical microscopy. In contrast to quadrant detection and related optical trap techniques, which present a substantial technical challenge for many users and provide low statistical power, DLS μ R is readily implemented on a commercial benchtop platform and delivers statistically averaged quantities with minimal user “hands-on” time.

We exploit these capabilities to interrogate the hierarchical molecular relaxations that occur in polymeric gels and precious biological materials, including DNA, extracellular matrix, and intestinal mucus. Our measurements capture physical processes ranging from the rapid bending fluctuations of individual polymers to the long-time-scale relaxations in entangled macromolecular networks, and are rationalized in terms of existing polymer physics theories. Together, these findings demonstrate that DLS μ R is a powerful tool for exploring the rheological behavior of a variety of precious soft and biological materials that can be readily adopted by an expansive scope of researchers.

METHODS

Detailed experimental methods are provided in the [Supporting Information](#).

ASSOCIATED CONTENT

Supporting Information

The Supporting Information is available free of charge on the [ACS Publications website](#) at DOI: [10.1021/acscentsci.7b00449](https://doi.org/10.1021/acscentsci.7b00449).

Figures, detailed experimental methods and analysis, a derivation for the predicted high-frequency scaling behavior of polyacrylamide gels, theoretical analysis of wormlike chains, comparison between microrheology and macrorheology for Matrigel, and DLS μ R experiments (PDF)

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B.A.K. designed research, conducted research, contributed new analytical tools, and wrote the paper. C.T. designed and conducted research and wrote the paper. P.D. and A.Z. conducted research. J.L.S. designed research. S.C.H. and A.J.S. designed research and wrote the paper.

Notes

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