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Transient grating spectroscopy nondestructively characterizes the mechanics of rubbery polymers and soft gels

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Designing efficient mechanisms for moving mechanical assemblies requires the use of materials with well-defined mechanical responses. Appropriate methods are needed to characterize these mechanical responses. Mechanical characterization of soft materials is critical in the high strain rate regime where intuition from manipulating a material at low rates fails to translate to applications including impact protection, tire traction, and sound damping. Here, transient grating spectroscopy (TGS) is used to measure wave propagation for soft elastomers and hydrogels. TGS is a non-destructive and non-contact optoacoustic technique that enables high strain rate measurements of the bulk modulus, a measure of a material's resistance to changing volume. The bulk modulus of elastomers and hydrogels is measured using TGS and its conversion to Young's modulus is discussed. This data is used to resolve values of Poisson's ratio in nearly incompressible gels to high degrees of precision.

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1. Introduction

Mechanics aims to control the dynamics of systems that provide useful action by understanding the storage, transfer, recovery, and dissipation of energy. Materials are the conduit through which this action occurs and are inherent sources of dissipated energy. Thus, characterizing and controlling the dynamic mechanical response of materials is key to developing efficient mechanisms. Nowhere is this characterization more important than in the high strain rate regime ($\dot{\epsilon} > 10^6$ Hz)¹ where materials, such as rubbery crosslinked polymers and soft gels, display behavior that is markedly different from their low strain rate response. This contrast means that any intuition researchers can glean from characterizing a soft material at low strain rates does not necessarily translate to the high strain rate regime. Accordingly, high strain rate characterization is vital to designing soft materials for impact protection,² sound damping,³ tire traction,¹ and the dynamic motion of small robots and organisms.^{4,5}

While the high strain rate response of soft materials is important in applications, characterizing this behavior is challenging. Common mechanical characterization techniques (e.g., dynamic mechanical analysis,^{6,7} shear rheometry,^{8–11} and

atomic force microscopy)^{9,12} that rely upon applying a deformation and measuring the stress response are fundamentally limited to timescales above the time it takes a wave to propagate across the sample to the other end and back. As shown in the SI, this wave speed limit can be estimated and is shown in Fig. 1. In practice, such measurements become unreliable at much lower frequencies in the kHz range where inertial effects overwhelm the signal from the material.^{13–15} Here, strain rates higher than the inertial limit line are only accessible through time-temperature superposition (TTS) methods, which tend not to work with gels due to evaporation and solvent ejection at rising temperatures. Fig. 1 highlights the divide between low and high strain rate characterization methods that may be bridged through TTS.

High strain rate techniques such as quartz crystal rheometry,²⁰ cavitation rheometry,^{16,21–23} and Split-Hopkinson (Kolsky) bar measurements²⁴ all overcome this limit by imposing deformation and then measuring the wave speed to quantify material behavior. These techniques have been very successful for high strain rate characterization yet have potential shortcomings such as the need for a thin film geometry, destructive test nature, and need to contact the sample, respectively. The recent development of transient grating spectroscopy (TGS) provides another pathway to high strain rate characterization that is not subject to these shortcomings.²⁵

Transient grating spectroscopy (TGS) is an optoacoustic technique that creates a temporary interference pattern within a sample by crossing two laser beams causing a periodic modulation in sample properties. Through probing the evolution

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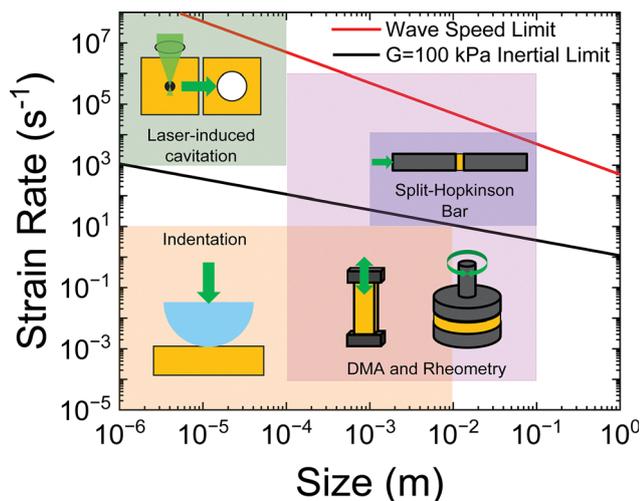


Fig. 1 Comparison of techniques and the relationship between sample size and achievable strain rates.^{16–19}

of this interference pattern over time, transient grating spectroscopy can provide insights into various dynamic processes, such as diffusion, energy transfer, and structural changes in materials. Its ability to capture ultrafast phenomena in real-time, from femto-second to microsecond timescales, has positioned TGS as a powerful tool for understanding fundamental processes in chemistry, physics, and materials science.^{25–28} Based on this, we aim to exploit TGS to explore the altered mechanical behavior of soft materials at high strain rates.

This paper addresses this goal by combining high strain rate TGS measurements with low strain rate indentation to characterize the elasticity of soft materials. The materials examined in this study range from rubbery crosslinked silicone elastomers to soft, biocompatible hydrogels. The use of low and high strain rate techniques provides complementary perspectives on the mechanical response of soft materials. Particular attention is paid towards exploring the manner in which these two perspectives can be related to one another. The results presented below have clear implications for predicting and controlling the mechanical behavior of soft materials in high strain rate applications.

2. Materials and methods

A. Polymer gels

A1. Polydimethylsiloxane (PDMS). Polydimethylsiloxane (PDMS) elastomer samples were prepared by combining Sylgard 184 silicone elastomer base with Sylgard 184 silicone elastomer curing agent at different ratios and curing temperatures. The oligomer-to-crosslinker ratio was varied between 10:1 and 20:1 by mass, and Rhodamine B dye was added to achieve a final concentration of 0.025 wt%. A speed mixer (FlackTek SpeedMixer DAC 150.1 FVZ-K) was used to evenly mix all constituents together at 2000 rpm for 40 s, and the mixture was dispensed into a polystyrene or glass dish and cured at room temperature for five days or at 120 °C for 3 h. Note that the sol fraction was not extracted and the samples

were tested in the as-formed state. The samples for indentation were prepared by punching out disks with a 16 mm diameter and ~3 mm thickness.

A2. Polyacrylamide (PAAm). Polyacrylamide (PAAm) hydrogel samples were prepared with varying monomer and crosslinker concentrations through free radical polymerization of acrylamide (AAM) monomer with *N,N'*-methylenebisacrylamide (MBAm) crosslinker. Ammonium persulfate (APS) initiator and *N,N,N',N'*-tetramethylethylenediamine (TEMED) catalyst were used to initiate the reaction. Stock solutions of AAM (30 wt%), MBAm (2 wt%), TEMED (10 vol%), and APS (10 wt%) were prepared in ultrapure deionized (DI) water (18.2 MΩ cm). Aliquots of each constituent were used to form a precursor solution with varying AAM (3.75, 7.5, 10, 12.5, 17.5 wt%) and corresponding MBAm concentration (0.15, 0.3, 0.4, 0.5, 0.7 wt%), following the methods by Uruena *et al.*²⁹ The monomer to crosslinker ratio was held constant, and the APS and TEMED concentrations were held at 0.15 wt% across all hydrogels. Rhodamine B dye was added to the precursor solution to achieve a final concentration of 0.025 wt%. The precursor solution was polymerized against polystyrene plates, and hydrogel sections (16 mm diameter, ~3.8 mm thickness) were equilibrated in ultrapure water at least 24 h before testing. After swelling, samples were ~4 mm in thickness.

A3. Poly(hydroxyethyl methacrylate) (PHEMA). Poly(hydroxyethyl methacrylate) (PHEMA) hydrogel samples were prepared by polymerizing hydroxyethyl methacrylate (67 wt%), MBAm (0.2 wt%), APS (0.15 wt%), and TEMED (0.15 wt%) in ultrapure water. The precursor solution was molded within a polystyrene dish and placed in the oven (Fisherbrand Isotemp Model 281A) at 50 °C for 2 h while under a nitrogen environment. After polymerization, the hydrogel samples were sectioned with a 16 mm diameter punch (~3 mm thickness) and equilibrated in solutions with varying water and ethanol volume fractions (0, 65 vol% and 100 vol% ethanol). The gels were swelled in Rhodamine B dye to ensure a final concentration of 0.025 wt%. Hydrogel samples were equilibrated for at least 24 h prior to experimentation.

B. Techniques

B1. Microindentation. Microindentation measurements were conducted using a custom microindenter (Fig. S1) to determine the reduced elastic modulus, E^* , of the samples. With an assumption of Poisson's ratio, $\nu = 0.5$, E^* can be converted to E with a factor of approximately 1.33 due to the relationship $E^* = E/(1 - \nu^2)$. The strain can be expressed by eqn (1),

$$\varepsilon \approx \frac{d}{R} \quad (1)$$

where R is the radius of curvature of the indenter probe and d , is the indentation depth. From this expression we can obtain a strain rate that uses the velocity of the indenter, v , and R to arrive at eqn (2), below.

$$\dot{\varepsilon} = \frac{v}{R} \quad (2)$$

The maximum confinement ratio observed in this work is $\frac{a}{h} = 0.25$ (a is contact radius and h is sample height) suggesting



that finite thickness effects should not significantly impact this measurement.

B2. Transient grating spectroscopy. TGS is a non-contact, non-destructive method of characterizing a material that uses optoacoustic signals to measure different properties.²⁵ It employs two different laser sources: a pulsed laser as a pump and a continuous wave laser as a probe as shown in Fig. 2a. Sample heating is minimized by only subjecting the sample to the lasers during the measurement time window. The power of each laser was also reduced to avoid any visible damage to the surface of the sample. To avoid dehydration, gels were reswollen between measurements. The pump laser employed here is elliptical in shape with a 5 mm major axis and a 0.5 mm minor axis. The continuous wave measurement beam has a diameter of 330 μm . At the phase mask, the pump beam is diffracted and a pair of first order diffracted beams are created. A phase mask consists of a transparent (*e.g.*, glass) substrate with a set of grating patterns with different periods etched on the substrate. The period of the optical grating formed on the sample surface can be easily adjusted by translating the phase mask between different etched grating patterns. The local heating generated by the pulsed beam through the grating induces a thermal or acoustic wave causing variations in stress or density. The probe beam is then diffracted off the variations caused by these waves and

the output signal (Fig. 2b) can be used to determine acoustic velocity, thermal diffusivity, and elastic properties.

Since the wavelength of the launched acoustic waves match the grating period L on the sample (which is half of the grating period on the phase mask based on optical parameters used in our system; see SI for more details), dispersion relations of the acoustic waves can be mapped by measuring the frequencies of the oscillations in the TGS signal at a range of grating periods (Fig. 2c). For the polymers studied here, a Fourier transform of each output was used to determine the frequency of oscillation. Note that the sharpness of these peaks indicates that the error in the measurement of oscillation frequencies is at largest 0.01 GHz (from the width of the peaks), which is an order of magnitude below the observed peak values. Depending on the stiffness of the material, the acoustic wave frequency detected by optical TGS is typically in the MHz to GHz range (Fig. 2d). The frequency of these acoustic waves can then be translated to a bulk modulus through eqn (3),

$$v = Lf = \sqrt{\frac{K}{\rho}} \quad (3)$$

where K is the bulk modulus and ρ is the density of the material. As can be seen in Fig. 2d, the linear relationship between frequency and inverse grating period suggests that the

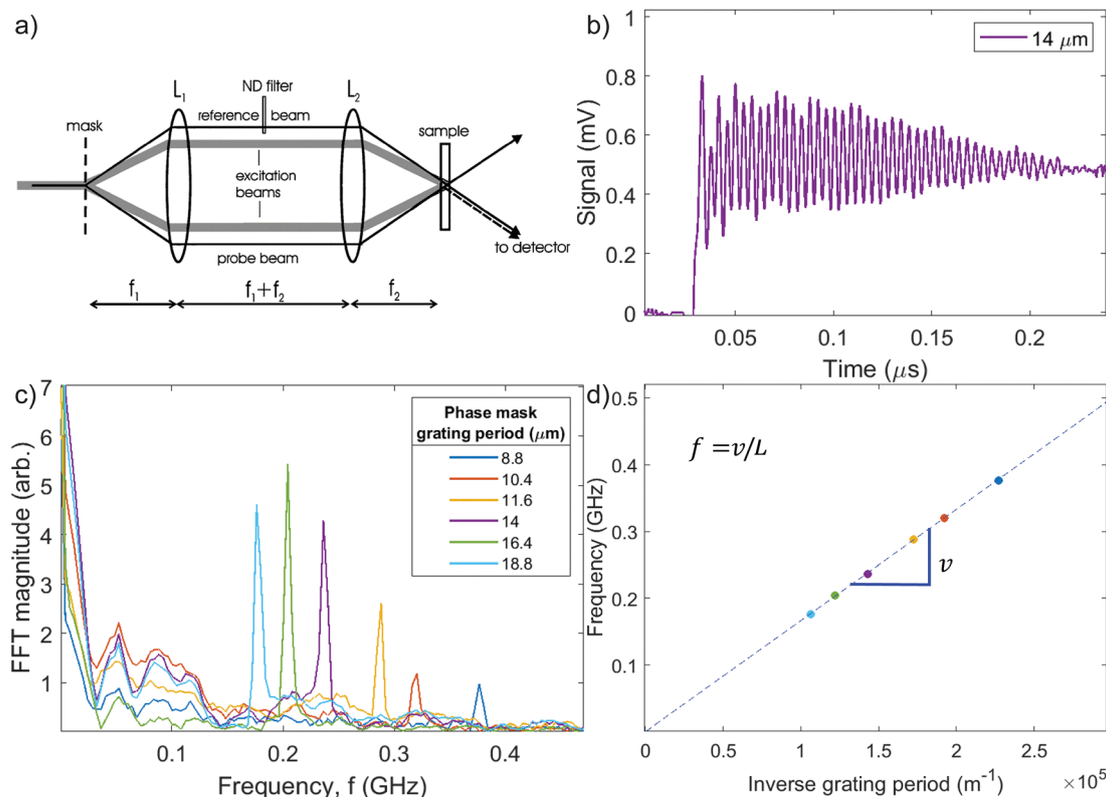


Fig. 2 (a) Schematic of TGS used for polymer measurements. Reproduced with permission from ref. 28. Polyacrylamide sample with 17.5 wt% AAm: (b) TGS trace, or signal, for one grating period. (c) Fourier transform of the resulting trace to determine frequencies of oscillation at each grating period. (d) Dispersion of grating period vs. frequency to find the bulk longitudinal wave speed. The wave speed for this sample of 17.5 wt% AAm was approximately $1571 \pm 3.4 \text{ m s}^{-1}$.



material wave speed is constant across these experimental conditions. The surface acoustic waves used in this measurement are sensitive to a depth of approximately $\frac{L}{2}$.²⁷ This relationship is helpful in characterizing the mechanical properties of a given material.

To estimate the strain rate, it is necessary to estimate the thermal strain during the measurements. The absorption coefficient of the Rhodamine B dye at 532 nm can be estimated as $10^5 \text{ M}^{-1} \text{ cm}^{-1}$. Combining this with the molarity of the dye in the gels, $5.2 \times 10^4 \text{ M}$, gives an absorption depth of approximately 0.02 cm. Assuming the pulse energy is absorbed within this depth, the temperature rise can be estimated from the pump laser fluence of 158 mJ cm^{-2} and the specific heat of water $4.2 \text{ J cm}^{-3} \text{ K}^{-1}$. Combining these values gives an estimated temperature rise of 2 K during the TGS measurement. This temperature rise can be combined with the thermal volume expansion coefficient of a hydrogel $2 \times 10^{-4} \text{ 1/K}$ to predict a volumetric thermal stretch of 1.0004. Assuming an isotropic expansion, the strain in one direction can be estimated by taking the cubic root of the stretch and subtracting 1 to get a strain value of 1×10^{-4} . The strain rate can then be estimated by taking this strain value and dividing by the 10^{-8} s each oscillation takes to get an estimated strain rate of 10 000 1/s.

3. Results and discussion

The output of TGS shows the time-domain signals that represent the intensity of light as a function of time (Fig. 2b). This is in the form of acoustic waves oscillations that are characteristic of the material behavior. In the case of one hydrogel sample, PAAm with 17.5 wt% AAM (Fig. 2), it is apparent both that the observed frequency of oscillation, f , can be simply extracted through a fast Fourier transform and that f varies as the grating period is altered. When this frequency is compared across all grating periods, a bulk acoustic wave speed can be determined from the slope in Fig. 2D. The values of each of the gels tested are shown below in Table 1. For PAAm, there is a slight, systematic increase in the observed wave speed as the initial polymer content increases. The wave speed values measured here for PAAm hydrogels span values from 1523 m s^{-1} to 1571 m s^{-1} and are consistent with the 1500–1650 m s^{-1} values measured by Gorman and McNeil through Brillouin Light Scattering (BLS).³⁰ For PDMS, the bulk acoustic wave speed is constant within the error of the measurement. For the stiffer formulations of PDMS, the variation is larger due to more scattered light, impacting the wave speed measurements of the samples. The agreement of values across PDMS samples suggests that the measured wave speed is insensitive to the crosslinking density and sol fraction. To ensure consistency across samples, the same power, as discussed in more detail in the SI, was used for the same grating periods as increasing the power can lead to heating that modifies the local properties of the spot being measured. The reported $1072 \pm 311 \text{ m s}^{-1}$ value measured here for PDMS agrees well with a reference value of 1319 m s^{-1} measured with BLS in the work of Stevens *et al.*³¹

Table 1 Wave speeds calculated using transient grating spectroscopy measurements. 3 measurements were performed at different locations on each sample and averages were calculated from the response of 5 different samples. The error values represent the standard deviation of the measured distribution across samples

Polymer	Bulk longitudinal wave speed (m s^{-1})
PAAm	
3.75 wt%	1523 ± 2.17
7.5 wt%	1551 ± 10.2
10 wt%	1561 ± 4.64
12.5 wt%	1570 ± 5.01
17.5 wt%	1571 ± 3.37
PDMS	
10:1–25 °C	1072 ± 311
10: –120 °C	1093 ± 306
20:1–25 °C	1062 ± 8.5
20: –120 °C	1073 ± 69.9
PHEMA	
Swelled in 100% ethanol	1225 ± 37.2
Swelled in 65% ethanol	1482 ± 95.1
Swelled in 0% ethanol	Not measurable

PHEMA shows a marked difference in wave speed for the gel as the solvent is altered to contain more water. Due to increased light scattering, the PHEMA sample in 100% water was not reliably measurable.

As is apparent from eqn (3), the wave speed can be converted to the bulk modulus when the density of the sample is known. This calculation has been undertaken and the moduli are plotted in Fig. 3 against variables relating to the formulation of each type of gel. The x -axes for the PAAm hydrogels, PDMS elastomers, and PHEMA gels are the initial concentration of reagents before network formation and swelling, weight ratio of base:curing agent, and the water content of the ethanol/water swelling solvent, respectively. E from indentation is shown in the first row and K from TGS is shown in the second row. For the PAAm hydrogels in Fig. 3a and d, it is apparent that the slight sensitivity of K to the initial polymer concentration is much less pronounced than that of E . This difference in sensitivity is likely due to the different physical origin of the two moduli. K results from the intermolecular forces that resist volumetric changes in rubbery polymer networks.³² In contrast, E in rubbery polymer networks results from the entropic penalty of stretching chains.³³ Since K is related to intermolecular forces, which should not change overly much for the highly swollen PAAm hydrogels (mainly composed of water), the relative insensitivity of K to initial polymer concentration makes sense. By similar reasoning, since E is related to the stretching of polymer chains, which is directly affected by the amount of polymer in the hydrogels, their sensitivity to the initial polymer volume fraction is reasonable.

In looking at the PDMS elastomers in Fig. 3b and e, it is apparent that E is sensitive to the cure temperature and crosslink density while K is not. Note that E is measured from indentation and is characterized at much lower strain rates than the TGS method used to characterize K . This is expected as K derives from intermolecular forces and is mainly sensitive to compositional variations instead of structure variations.³²



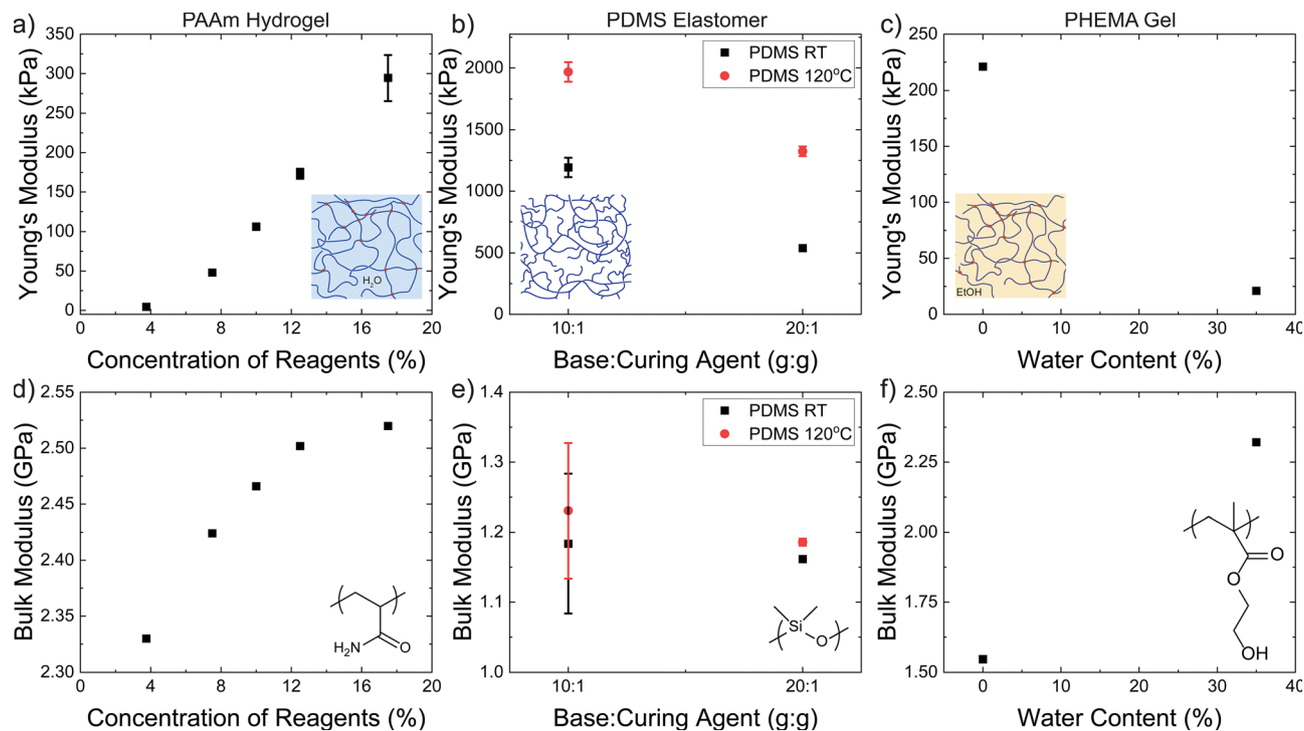


Fig. 3 Comparison of (a–c) Young's modulus and (d–f) bulk modulus measurements of (a, d) PAAm hydrogels, (b, e) PDMS elastomer, and (c, f) PHEMA gels. The strain rate for the calculation of the Young's modulus is ~ 3 Hz and the strain rates for the bulk modulus are on the order of 10^4 Hz. The observed trends are consistent with the different physical origin of the two moduli.

The main difference in the PDMS elastomers at different mixing ratios and cure temperatures is in structure, namely in the network crosslinking density and sol fractions, while the samples are all composed of the same silicone molecules. The agreement between values is consistent with the understanding that, while E is sensitive to the structural features of the network, K is only sensitive to the intermolecular interactions present in the material which are unaltered by changing the crosslinking density and sol fraction. This results in significant changes in E despite a constant observed value of K . The PHEMA gels in Fig. 3c and f are subject to additional compositional effects. Here both E and K show marked differences in value with respect to the swelling solvent composition. E shifts in this case because using mixed solvents alters the solvent quality and changes the equilibrium polymer volume fraction after swelling. In addition to changing the solvent quality, mixing solvents alters the intermolecular forces that determine K . Ethanol has a lower K than water which explains the decrease in K observed in Fig. 3f when the gel is swollen in 100% ethanol.

Notably, the observed K value for the PDMS elastomers is less than that for both the PAAm hydrogels and the PHEMA gels despite the fact that E is an order of magnitude greater in the PDMS elastomers. This difference shows how the mechanical intuition that researchers can gain by manipulating a soft solid by hand does not necessarily translate to the high strain rate regime. In other words, when designing for high strain rate applications, it is critical that researchers make decisions informed by high strain rate measurements.

While the comparisons between moduli and structural parameters presented above makes qualitative sense, each of these moduli are measured at extremely different strain rates. The indentation measurements were performed at strain rates of approximately 3 Hz while TGS probed the materials at 10^4 Hz. Bridging these two strain rate regimes to connect these moduli measurements would be very impactful to our understanding of design in high strain rate applications. An initial attempt at doing so by compiling literature data collected at various frequencies is contained in Fig. 4. This figure shows E values for the PAAm hydrogels and PDMS elastomers measured at increasing frequencies. As is apparent in Fig. 4a, data for the PAAm hydrogel is relatively sparse. This is likely due to the ill-suited nature of PAAm hydrogels to the application of a TTS analysis. Specifically, shifting temperature is difficult in hydrogels swollen to equilibrium because doing so will both shift the swelling equilibrium and lead to an increase in sample drying rates.³⁴ The only high strain rate data on PAAm hydrogels is from laser-induced cavitation and is in the GPa range compared to the 10–100 kPa values observed at low strain rates.¹⁶ For the PDMS elastomers, TTS can be applied and thus more data is available; however, as a commercial product whose final properties depend heavily on mixing and curing procedure, data compiled in this manner should not be considered authoritative. The trends for this system show an increase in E with frequency for the PDMS elastomers. For both materials, measurements of E do not exceed 10^7 Hz which is close to the frequency range accessible by TGS.



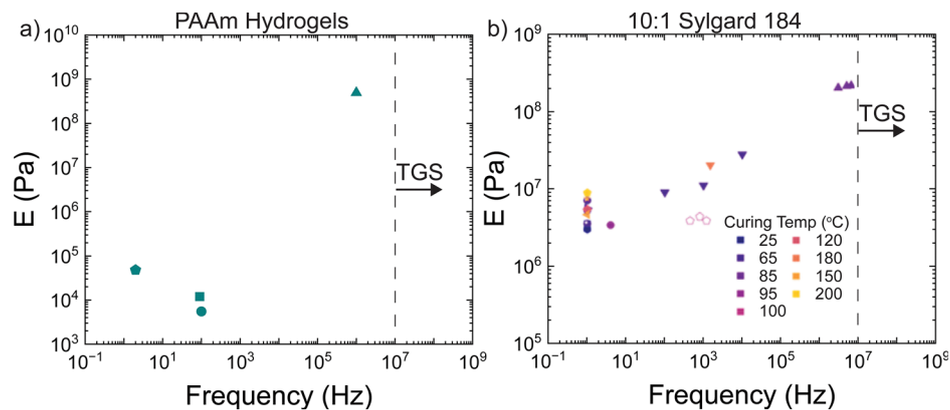


Fig. 4 (a) Frequency-dependent storage moduli in PAAm gels with 6–8 wt% AAm (square,³⁵ circle,⁸ triangle,¹⁰ pentagon: indentation). (b) Frequency-dependent Young's modulus for different curing temperatures of PDMS (Sylgard 184) in a 10:1 monomer: crosslinker ratio. The colors map the temperatures while symbols indicate sources. (circle,³⁶ right triangle,³⁷ up triangle,³⁸ left triangle,³⁹ down triangle,⁴⁰ filled pentagon,⁴¹ pentagon,⁴¹ diamond⁴²). These values may be more qualitatively compared since there are a combination of techniques that calculate modulus values in different ways. The target range that TGS measures is indicated on each plot.

The gap in available data on E in the high strain rate regime is interesting as TGS can potentially measure this quantity in that regime. As illustrated in Fig. 5, the only real hurdle in quantifying E in the high strain rate regime is the need to convert measurements of K to E . From the mechanics perspective, this is a problem with a straightforward solution where $E = 3K(1 - 2\nu)$ and ν is Poisson's ratio.⁴³ Note that the relationship between these quantities is derived using a linear elastic constitutive relationship. Thus, the problem becomes a question of having a high strain rate value of ν . Unfortunately, to the best of the authors' knowledge, measurements of ν in these nearly incompressible systems have been limited to quasi-static measurements. While K is relatively insensitive to strain rate when compared to E ,⁴⁴ ν is sensitive to strain rate due to its dependence on E . Since ν varies at different strain rates this precludes the conversion of K to E when ν is not quantified at the same strain rate as K . Thus, high strain rate measurements of ν are needed to extract high strain rate values of E from TGS.

One potentially interesting result that comes from considering the interconversion of these moduli is that while the conversion between E and K is sensitive to the value of ν used, the calculation of ν from E and K is largely insensitive to errors

in these two moduli. The experimental error in calculating ν from E and K is given by eqn (4),

$$\Delta\nu = \frac{E}{6K} \sqrt{\left(\frac{\Delta E}{E}\right)^2 + \left(\frac{\Delta K}{K}\right)^2} \quad (4)$$

where it is apparent that any error in either modulus ends up being scaled by the ratio of these two moduli.⁴³ These quantities are typically at least 3 orders of magnitude apart so large errors in the moduli values can still produce accurate values of ν . For example, if $E = 1$ MPa and $K = 1$ GPa and each modulus has a $\pm 25\%$ error this propagates to a 0.01% uncertainty in the calculation of ν meaning that ν can be reported out to 4 decimal places. Calculating ν still faces the fundamental problem that the two moduli value are gathered at extremely different strain rates. However, K is relatively insensitive to changes in frequency compared to E and ν is insensitive to errors in K .⁴⁴ This means that any potential increase in K in the high strain rate regime could, in this one very specific instance, be treated similarly to an effective measurement error for K values in the low strain rate regime. For example, the 10:1–120 °C PDMS elastomer samples have a value of $K = 1.4$ GPa from TGS

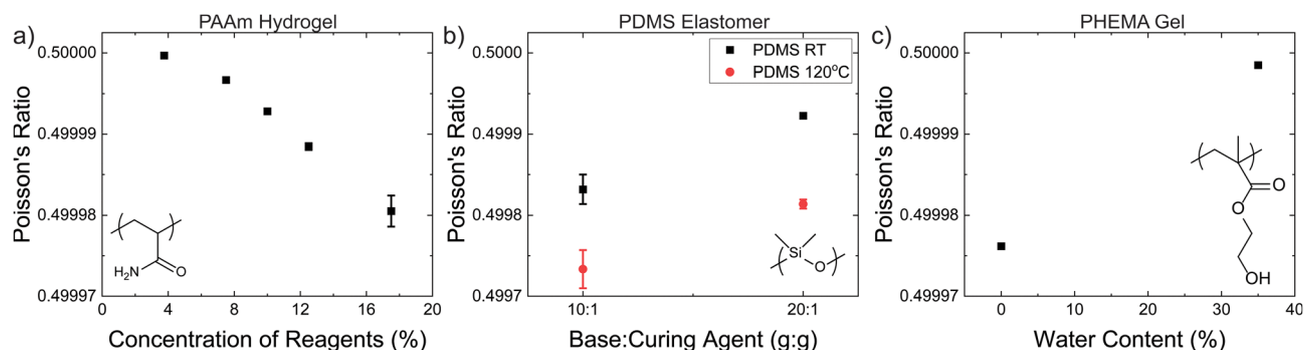


Fig. 5 Calculation of Poisson's ratio using values from indentation and TGS measurements for (a) PAAm, (b) PDMS, and (c) PHEMA. Within the narrow range of potential values of ν , the results follow the expected inverse trend with respect to the Young's modulus.



performed at 10^4 Hz while indentation gives $E = 2$ MPa at about 3 Hz. When combined these moduli give a calculated ν value of 0.49976 which is close to a quasi-static reference value of 0.4998.⁴³ This suggests that high strain rate values of K can be combined with low strain rate values of E to quantify low strain rate values of ν .

While the conversion of K to E is stymied by the lack of available literature data on high strain rate ν , it is clear that TGS can be used to estimate low strain rate values of ν . Values from this analysis have been calculated and are shown in Fig. 5 for the different gels plotted against variables relating to the formulation of the gels. For the PAAm samples, ν decreases with increasing E values which increase at higher initial concentration of reagents. The values for PDMS decrease in stiffer samples also indicating an inverse relationship with E . A similar trend is observed for PHEMA where ν decreases in gels swollen with pure ethanol when E is highest. Notably, the values of ν for PAAm hydrogels are significantly higher than those previously measured in the literature by applying optical strain gauge analysis to uniaxial extension measurements.^{45,46} The confidence in the measurements presented in this work derive from arguments quantifying the propagation of errors and the sensitivity to the presence of cumulative errors.⁴⁷ As presented above in the discussion of eqn (4), the propagation of errors shows that incredibly precise values of ν can be calculated from combining E and K when they are at least three orders of magnitude apart. Beyond precision, the accuracy of the mean reported values of ν is sensitive to the presence of cumulative errors that propagate from the individual measurements of E and K . The sensitivity to such errors was recently quantified by Nedoluha *et al.*⁴⁷ and it was found that, of the four ν calculation methods analyzed, inferring ν from E and K was the method with the lowest sensitivity to cumulative errors. These results highlight the strength of TGS as an accurate and precise method to characterize the Poisson's ratio of nearly incompressible gels and elastomers.

While TGS was ultimately unable to estimate high strain rate values of E without a high strain rate value of ν , TGS still provides value to the soft materials community by characterizing low strain rate values of ν in difficult to measure samples. Note that the combination of low strain rate and high strain rate data used to calculate low strain rate ν values in this work is only appropriate due to both the relative insensitivity of K to strain rate when compared to E and the insensitivity of ν to errors in E and K measurements when the two moduli are at least three orders of magnitude apart. Future researchers should be cautious when attempting to combine data measured at different strain rates as is done in this work. Developing high strain rate testing methods capable of quantifying Poisson's ratio remains an open question. TGS is advantageous in this space as it provides a non-contact, non-destructive means of quantifying K for hydrogels that are difficult to manipulate in other techniques. This capability unlocks current and future measurements of ν in soft gels where existing methods lack the resolution to meaningfully distinguish nearly incompressible values from the incompressible limit.

4. Conclusion

Transient grating spectroscopy is a nondestructive and noncontact technique that has potential to measure high frequency responses of soft materials, particularly in the context of hydrogels like polyacrylamide (PAAm) and elastomers such as polydimethylsiloxane (PDMS), both of which are commonly used and generally well-characterized. While the output can directly be used to calculate the bulk acoustic wave speed and the bulk modulus for polymers, there is a lack of information about Poisson's ratio (ν) at higher frequencies that can bridge the gap to finding dynamic Young's modulus values. Here, in conjunction with a quasistatic method such as indentation, a low strain rate ν can be reliably calculated which provides more information on quasistatic properties. While the current body of research has primarily focused on Poisson's ratio values at lower frequencies, there exists a promising avenue for future studies to expand our understanding of higher frequency regimes by focusing on analysis of high frequency values of ν . Knowledge of dynamic Poisson's ratio values will lead to enhanced precision in characterizing material behavior under various conditions. In turn, this information can pave the way for the development of innovative materials and structures that can withstand and adapt to dynamic and high-frequency or high strain rate mechanical environments. This study also underscores how transient grating spectroscopy holds significant promise for being a dynamic test platform of polymers provided they meet the constraints of the technique.

Conflicts of interest

The authors declare no conflicts of interest.

Data availability

The data supporting this article have been included as part of the supplementary information (SI). Supplementary information includes further details on the indentation and TGS methods, estimates of polymer mesh sizes, a reporting of the densities used during calculations, and the arguments used to estimate the limits in Fig. 1. See DOI: <https://doi.org/10.1039/d5sm00835b>.

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