Relationships among structure, memory, and flow in sheared disordered materials

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Abstract

A fundamental challenge for disordered solids is predicting macroscopic yield from the microscopic arrangements of constituent particles. Yield is accompanied by a sudden and large increase in energy dissipation due to the onset of plastic rearrangements. This suggests that one path to understanding bulk rheology is to map particle configurations to their mode of deformation. Here, we perform laboratory experiments and numerical simulations that are designed to do just that: 2D dense colloidal systems are subjected to oscillatory shear, and particle trajectories and bulk rheology are measured. We quantify particle microstructure using excess entropy. Results reveal a direct relation between excess entropy and energy dissipation, that is insensitive to the nature of interactions among particles. We use this relation to build a physically-informed model that connects rheology to microstructure. Our findings suggest a framework for tailoring the rheological response of disordered materials by tuning microstructural properties.

1 Introduction

Disordered solids are ubiquitous. They are found, for example, in our foods as pastes and gels\textsuperscript{1}, and amidst our homes in the form of concrete\textsuperscript{2} and mud\textsuperscript{3,4}. Frustratingly, these materials can experience sudden mechanical failure, such as the collapse of soil during rapid mudslides. Indeed, when sufficiently stressed, all disordered materials exhibit a swift decrease in ability to support load. In the vicinity of this “yield” transition, the solid material shifts from a state wherein energy is stored via internal elastic forces, to a state in which energy is dissipated via irreversible plastic rearrangements\textsuperscript{5–9}. Microscopic spatiotemporal features are associated with this yield transition and affect macroscopic material responses such as ductile versus brittle behavior. Unfortunately, in contrast to the case for crystalline materials, our ability to predict and control yield in disordered solids based on their constituents and their interactions is still limited\textsuperscript{6–10}. To build such microstructural models, we need to identify key microscopic metrics relevant to plasticity in disordered materials\textsuperscript{11}. Recently, excess entropy has been explored for this purpose\textsuperscript{12–14}. In equilibrium systems, excess entropy has been utilized to connect viscosity with interparticle structure\textsuperscript{15–18}. Recently in far-from-equilibrium systems, excess entropy scaling has been shown to facilitate a relationship between microscopic structure and dynamics\textsuperscript{12–14}. Thus, excess entropy offers an untapped signature for plasticity and a potential tool for modeling the mechanical response of disordered solids.
The study of rheology and particle dynamics in disordered systems has a venerable history. As a result of this research, theories have proliferated in recent decades. Two of the most successful are Mode Coupling Theory, wherein the interplay of dynamical modes causes the emergence of rearrangements, and Shear Transformation Zone theory, which posits that local configurations determine where rearrangements occur. More recently, structural signatures for rearrangement have been revealed by machine learning approaches, by study of low-frequency excitations, and via local yield stress and near-neighbor cage dynamics. Despite their usefulness, difficulties remain in applying these theories to experiments because of the need for fitting parameters and the use of empirical relations that are difficult to measure. Moreover, these theories typically do not account for history-dependent behavior such as material memory, which is necessary to understand plasticity.

Generally, disordered materials contain memories, i.e., microscopic signatures related to how the material has been processed. Memory of a previous shearing direction, for example, can be encoded into a material’s response; once a material is sheared sufficiently in a given direction, continued shear in that direction requires more force than in the opposite direction. In jammed systems, recent experiments and simulations have studied formation of directional memory at low strain amplitudes, both below and near the yield transition; far above yield, memories are erased. These observations, in turn, raise important new questions: do memories require elastic storage? Is plasticity synonymous with erasure? How do these phenomena manifest during yield, e.g., in storage and loss moduli?

In this contribution, we utilize excess entropy to quantify material memory and construct a microstructural model for disordered-material response and energy dissipation. Experiments and simulations show that three non-dimensional parameters govern the connections between microstructure and bulk rheology: packing density, a normalized (non-dimensional) form of the imposed stress, and an excess entropy (microstructure-related) ratio that quantifies the material’s ability to retain information about its initial state. Our results confirm that memory is stored elastically and lost plastically, and show how yield and the ductile/brittle response emerge from knowledge about particle configurations at the microscopic scale.

2 Results

The experiments investigate disordered solids. The solids are colloidal monolayers of athermal, spherical particles adsorbed at an oil-water interface. The charged particle surfaces generate a dipole-dipole repulsion between particles. This repulsion is strong enough to jam the entire material, arresting particle motions. To probe the effects of disorder, we study both mono-disperse and bi-disperse spherical particle systems with diameters of 5.6 μm and 4.1 μm-5.6 μm, respectively. In the bi-disperse system, crystalline domains tend to be much smaller (See Supplementary Materials). We impose many cycles of sinusoidal stress on these samples using a custom-made interfacial stress rheometer that permits measurement of the bulk response of the colloidal monolayer while simultaneously recording trajectories of individual particles (see Methods). Cyclic stress is quasi-static, insofar as the time scale for a completion of a rearrangement (∼0.5s) is much shorter than the shortest driving period (5s) or largest inverse strain rate (20s).

We investigate particle rearrangements by identifying non-affine deformations within each particle’s neighborhood. The degree of non-affinity is quantified by the mean-squared displacement after subtracting the best fit affine transformation, $D_{\text{min}}^2$ (see references for more information). Within cyclically sheared disordered materials, two types of non-affine events occur (Fig. 1a): those wherein particles return to their original position at the end of a strain cycle but along different paths, and those wherein particles escape their nearest neighbors and do not return. For visualization we define $D_{\text{min},C}^2 \equiv \pm \sqrt{(D_{\text{min},R}^2)^2 + (D_{\text{min},E}^2)^2}$, where the subscripts refer to returning (R) and escaping (E) events, respectively, and sign corresponds to the greater $D_{\text{min}}^2$. Both types of events dissipate energy. Returning non-affine events are known to emerge near the yield point when elasticity begins to diminish and plasticity starts to increase; escaping events arise well beyond yield (Fig. 1b). The fraction of particles undergoing non-affine events is $f_d$. By following the rearrangements, we develop understanding about trajectory dynamics within the microstructure, and we take steps towards our ultimate goal to relate microstructure to rheology.

To quantify structure, we characterize the inter-particle forces and particle configurations using the radial distribution function, $g(r)$. Since the material is jammed, the motion of each particle is arrested by its neighbors. This caging, and escape thereof, provides another lens for the non-affine motions mentioned above; when enough particles pass each other via small changes in the structure of their surrounding cage, the material yields. For quantitative
Figure 1: **Overview of structure, dynamics, and response.** We characterize the disordered solid bulk response to cyclic stress from evolving configurations of individual constituent particles. (a) Image of ~40,000 particles. Part of the raw image is shown (left). The scale bar is 200µm. Detected particle positions are also shown (right). For illustration, color indicates $D_{\min,C}^2$, which quantifies the degree to which a particle has followed a non-affine returning trajectory (blue), or a non-affine escaping trajectory (red). The particles in this image are experiencing yield ($\gamma_0 \sim 15.7\%$). (b) Quantification of the fractions of escaping and returning events versus total strain amplitude. Returning events rapidly increase near the yield point ($\gamma_0 \sim 3.0\%$). (c) The number of particles, $Z(r)$ within a radius, $r$ of a reference particle. The radius is expressed in units of $a$, the average distance between neighboring particles. Vertical dashed lines indicate the limit of the first shell of neighboring particles. Inset: radial distribution function, $g(r)$. (d) The measured strain of the material versus the imposed stress throughout a cycle. Both stress and strain are averaged stroboscopically over 25 cycles. The different ellipses correspond to separate runs at different imposed stress amplitudes. Here, the area enclosed is a result of the lag between stress and strain, which in turn quantifies the energy dissipated from the material.

Analysis, we compute $F^*$, the sum of the magnitudes of inter-particle forces acting on the average particle. Specifically: $F^* = 2\pi \rho \int_{0}^{r_N} \left(-\frac{\partial u}{\partial r}\right) g(r) r dr$; here $\rho$ is the number density of particles, $r_N$ is an upper cutoff distance below which nearest neighbor particles are found, $u$ is the pair potential function between any two particles, $\frac{\partial u}{\partial r}$ is the force acting between any two particles, and $g(r)$ is the sample radial distribution function as a function of separation $r$ (Fig. 1c; Methods). To determine $r_N$, we use the coordination number as a function of radial distance, $Z(r)$ (Fig. 1c). $Z(r)$ is derived from $g(r)$ and has been studied to characterize particle interactions and their effect on bulk materials. In our systems, neighbor shells are well defined by broad peaks in $g(r)$ separated by troughs (Fig. 1c-inset). The extent of the nearest neighbor shell is defined as the radius at which $Z(r)$ begins to increase rapidly for a second time (Fig. 1c-main).

We quantify disorder using excess entropy, the difference between the system’s entropy and that of its ideal gas...
Figure 2: Memory within microstructure. Microstructural anisotropy reveals signatures of memory. Below yield, anisotropic orientation remains unchanged regardless of shear direction. Orientation quantifies stored memory. Above yield, anisotropic orientation reverses freely to match the direction of shear, indicating a loss of memory. (a) Radial distribution function, \( g(x,y,t) \) at a time corresponding to one quarter of the way through a shearing cycle. We fit an ellipse to the first neighbor ring. This ellipse stretches and reorients over time indicating changes in structural anisotropy of the sample. Two elliptic fits are shown at two times, \( t=1.25 \) (---) and \( 1.75 \) (---). (b) Orientation of the sample microstructure over time as a function of strain amplitude. With increasing strain amplitude, the microstructure reorients to match the stretching axis. It first reorients completely at the yield point (3.2%). (c) Elongation quantified by the ratio of ellipse major and minor axis lengths \( (m/n) \) over time. Below yield, elongation oscillates directly with the strain; above yield, elongation oscillates with twice the frequency of strain perturbation. In b & c data are averaged stroboscopically over 25 cycles.
Microstructure elongates only once per cycle at $\omega$.

Microstructural anisotropy reveals a memory of the last direction the material was sheared above yield (Fig. 2). To remove internal stresses, each of our experiments is pre-sheared well above yield ($\gamma_0 \sim 50\%$); nevertheless, this protocol imprints an anisotropy into the sample set by the last shear direction. Previously it was shown that this type of material memory is imprinted into $g(x, y)^{44,45,62}$. Here, we find that this memory imprint is associated with the principal directions of shear (Fig. 2). Once a memory is stored, the memory is retained as long as the material is sheared elastically. Precisely when the material yields, all memory is lost, and the microstructure freely switches between both orientations. Taken together, these results indicate that materials store and express memories in the elastic regime but lose them in the plastic regime. Furthermore, recently we showed that orientational memory is stored most strongly within crystalline domains wherein particle rearrangements are most intensely suppressed$^{46}$.

![Figure 3: Entropy and material memories.](image-url)

Figure 3: **Entropy and material memories.** Variation of entropy provides means for predicting system response to a given strain amplitude. (a) Excess entropy, with the mean value subtracted, follows a sinusoidal response. Below yield, its oscillation frequency is the shear cycle frequency. At yield, the excess entropy signal has components at both the driving frequency and twice the driving frequency: the material is beginning to forget its initial state. Above yield, the entropy response oscillates almost exclusively at twice the shear cycle frequency. Black dots indicate experimental data. Red lines are fits to equation 1 with $T$ as the only fitting parameter. The experimental data are averaged stroboscopically over 25 cycles. (b) Amplitudes associated with the first and second harmonics are present within the $s_2$ signals. Note, that the second and first harmonic amplitudes cross each other at the yield point, $\gamma_0 = 3\%$, designated by the vertical dashed line (--.--).

We now use excess entropy to characterize and relate observations about imprinted memory to the system microstructure. Above yield, we find that structural response is independent of the direction of shear (Fig. 3a, $\gamma_0 = 6.8\%$); when the material is sheared in either direction, the excess entropy increases and decreases as the shear is reversed. Ostensibly, the material cannot sustain a memory above yield, because it is continually forced out of meta-stable states within the energy landscape. Near yield, however, the direction of shear has an effect on structural response (Fig. 3a, $\gamma_0 = 2.2\%$). Notice, $s_2$ does not increase as the material is sheared over the second half of a sinusoidal shear cycle. Finally, below yield, the direction of shear is important; shear in one direction produces
an increase in excess entropy, and shear in the other direction produces a decrease (Fig. 3a, $\gamma_0 = 0.7\%$).

As seen in figure 3b, the $s_2$ signals are sinusoidal. The first harmonic ($\omega$) decays and the second harmonic (2$\omega$) grows with increasing strain amplitude. The first harmonic is dominant below yield, and the second is dominant above yield. Therefore, the amplitude of the first harmonic of $s_2$ provides quantification of a stored memory, and the amplitude of the second harmonic characterizes the degree to which memory of the initial state is lost. Notice, these first and second harmonic amplitudes cross each other near the yield point.

To build a relationship between excess entropy and bulk rheology, we next investigate the connection of $s_2$ to the other dynamical metrics. For this comparison, we compute the ratio of the second to first harmonic amplitude, which we denote as $s_{2,h}$. We can relate $s_{2,h}$ to several quantities in our system (Fig. 4). For example, $s_{2,h}$ scales with the product of $F^*/F_0$ and $f_d$ (Fig. 4a), where $F_0$ is the amplitude of the prescribed shear force. This relationship between dimensionless parameters suggests that when the imposed force on the system grows larger than $F^*$, the microstructure begins to permanently change, losing stored memory. Rapid variation of $f_d$ also signifies the transition.

These findings build on recent work that links excess entropy and non-affine particle dynamics\textsuperscript{12,13}. Note that the scaling in the present case is quadratic because $f_d$ varies nearly linearly with the imposed force, $F_0$ (see Supplemental Materials). Finally, we find that the product of $s_{2,h}^2$ and $F_0/F^*$ scales linearly with $G''/G'$ (Fig. 4c). The scaling factor for this linear relationship is $2\phi/\pi^2$; here $\phi = \pi Na^2/A$ quantifies the particle spatial density, $a$ is the average nearest neighbor distance derived from the first peak of $g(r)$ (Fig. 1c: inset), and $A$ is the total area of the observed sample or simulation.

The yield phenomenology shown in Fig. 4c depends on four dimensionless parameters: $F_0/F^*$, $s_{2,h}$, $G''/G'$, $\phi = \pi Na^2/A$, and $a$ is the average nearest neighbor distance derived from the first peak of $g(r)$.

Figure 4: Comparisons of imposed force, microstructural excess entropy, and bulk rheology. a) The imposed force amplitude, $F_0$, normalized by the elastic force capacity, $F^*$, is plotted versus the excess entropy harmonic ratio, $s_{2,h}$ (in both mono-disperse and bi-disperse experiments). A fit of the data suggests a parabolic relationship (p-value=3.14x10$^{-13}$, and $r^2$:0.989), corroborating equation 2. b) The increase in the ratio of loss and storage moduli, $(G''/G')$ versus strain amplitude in both the mono-disperse and bi-disperse experiments (same legend for mono-disperse and bi-disperse experiments as panel a). Yield is signaled by the rapid increase in parameter values at about 0.03 strain amplitude. Inset: data from simulations employing Hertzian and Lennard-Jones interaction potentials. In both cases, markers are measured values and lines are predictions of equation 3. c) Left and right hand sides of equation 3. Notably, all parameters are measured. The solid diagonal line (—, slope of 1.0) represents equation 3. The slope of the best fit to the data is 0.981, p-value=4.43x10$^{-26}$, and $r^2$:0.944.
and the packing density \( \phi \). The ratio \( F_0/F^* \) characterizes the shear force exerted on the material relative to
the force required to cause rearrangements; when \( F_0/F^* \geq 1 \) plasticity is non-negligible. The microstructural
quantity \( s_{2,h} \) provides a metric for whether a material’s response is dominated (or not) by memory as it experiences
oscillatory strain; this microstructural property can be interpreted as the degree of plastic strain. Finally, a
familiar ratio quantifies the bulk rheological response of the material: \( (G''/G^*) \). All experimental (and simulation)
data are collapsed using these dimensionless parameters, and a direct relationship between rheology, dynamics, and
microstructure is experimentally established in the disordered solid.

Numerical simulations complement the experiments. The simulations enable us to vary features of the disordered
system that are difficult to control experimentally. In particular, we can test ideas regarding variation of inter-particle
potential. Moreover, unlike the experimental system, which involves a fluid-fluid interface that gives rise to viscous
drag on the particles, the simulations offer the possibility to study the validity of our new concepts in disordered
materials without viscous drag. Thus, we have conducted shear simulations without viscous drag and with two
different inter-particle interaction potentials: Lennard-Jones, a model for atomic glass, and Hertzian, a model for
granular systems (see Methods).

The simulations and experiments exhibit remarkably similar behaviors. Across both the experiments and sim-
ulations, a direct and common functional relationship between excess entropy and rheology is revealed (Fig. 4c). This relationship does not depend on the details of particle interactions, nor the amount of disorder. Further, since
simulations do not involve a background fluid, the importance of hydrodynamic effects is ruled out. To test the limits
of applicability of our numerical findings, we introduce varying amounts of Brownian motion into the Lennard-Jones
simulations. At high thermal temperature, the particles rearrange due to Brownian motion in addition to shear
stress, and memory cannot be formed. However, at low thermal temperature, the experimentally observed relation-
ship between entropy and rheology holds (see Fig. 4b&c). Moreover, we find that jamming is required for the storage
of memories in both simulation systems. At low packing densities, where the system easily un-jams during shear, the
relation is violated. The wide applicability of these ideas suggests the existence of a deeper theoretical formulation.
Thus, in the remainder of this paper we outline how our results may be derived phenomenologically (for the full
derivation see the Supplemental Materials).

To elucidate the relationship between \( s_2 \) and the material properties \( (G', G'') \), we perform a simple energy balance. We start with the harmonic behavior in \( s_2 \). In this situation, energy is balanced in terms of accumulation, \( T \Delta S_2 \),
reversible (quasi-static) energy transfer, \( F^* x/2 \), and irreversible dissipation, \( f_d F x \):

\[
T \Delta S_2(t) = F^* x(t)/2 + f_d F(t) x(t).
\]

(1)

Here \( x(t) \) is the displacement of the system boundary, \( F(t) \) is the imposed shear force, and \( T \) is a parameter (generally
different from the thermal temperature) that converts differences in entropy to differences in energy.\textsuperscript{14,65–67} Note
that this equation would not apply in a system dominated by thermal motion, because we do not account for changes
in entropy due to thermal fluctuations. The equation also implicitly reflects the requirement of jamming via the \( F^* \)
term. With a single fitting parameter, \( T \), the changes in harmonic behavior in excess entropy are reproduced from
below to above yield (Fig. 3a).

The harmonic transition, associated with the excess entropy found in experiments and simulations, is captured
by the first and second terms on the right-hand-side of equation 1. \( s_{2,h} \) is the ratio of those two terms:

\[
s_{2,h}^2 = f_d F_0/F^* .
\]

(2)

This relation describes the harmonics data remarkably well (Fig. 4a). We next build on equation 2 by incorporating
a finding of shear transformation zone theory, namely that elastic energy builds up in the microstructure until
it is plastically released via non-affine rearrangement events\textsuperscript{22,23}. Quantitatively, this concept is represented as:

\[
G'' \propto N f_d G', \quad \text{where } N \text{ is the number of total particles observed; when substituted into Eq. 2 we obtain:}
\]

\[
\frac{G''}{G'} = \frac{2 \phi}{\pi^2} \frac{F^*}{F_0} s_{2,h}^2 .
\]

(3)

Note, that each parameter in this expression is measured and is generally accessible in many systems. Across
strain amplitudes, remarkable agreement is found between \( G''/G' \) measured in experiments and simulations, and the
predictions by Eq. 3 (see Fig. 4b&c).
3 Conclusion

Our results demonstrate that the yield transition of jammed systems has a configurational origin rooted in the persistence of material memory. We investigated the responses of several jammed systems undergoing cyclic shear deformation, incorporating aspects of STZ theory, excess entropy, and harmonic analysis into a single framework. The analysis reveals two new dimensionless parameters and three relations, derived phenomenologically, which connect particle configurations to bulk rheology. Importantly, the microstructural information needed, i.e., the radial distribution function, is available in myriad of scattering/microscopy experiments spanning length scales and particle types; thus, this analysis is accessible to experimentalists. In the future, it should be interesting to search for similar relations for other loading conditions, such as compression or steady shear, and to explore a wider array of particulate systems in which the particles are not simple spheres.

We have developed a framework to understand bulk properties of jammed materials under shear based on microstructural information. The findings hold potential to predict behavior of a broad range of dynamically arrested disordered materials including foams, gels, packings of nano- and micro-scale particles, and atomic/molecular glassy matter. Our findings, perhaps, also shed light on some deeper questions: in particular, the nature of entropy and the potential to use entropy ideas in far-from-equilibrium media. While entropy formulations for non-thermal systems have found utility in modeling disparate phenomena, its physical interpretation often remains mysterious. Disordered particulate packings appear to be particularly useful for clarifying this phenomenology, since their material structure can be interrogated with relatively simple methods.

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4 Methods

| ID | Type      | Forces   | Dispersity | Diameters | \(\phi\) | \(\Phi[\%]\) |
|----|-----------|----------|------------|-----------|--------|---------|
| A  | Experiments | Dipole-dipole | Bi-disperse | 4.1, 5.6\(\mu m\) | 14.02 | \(\sim\)31 |
| B  | Experiments | Dipole-dipole | Mono-disperse | 5.6\(\mu m\) | 13.99 | \(\sim\)35 |
| C  | Simulations | Lennard-Jones | Bi-disperse | N/A | 5.03 | N/A |
| D  | Simulations | Hertzian | Bi-disperse | 0.84, 1.16 | 9.68 | 110 |
| E  | Simulations | Hertzian | Bi-disperse | 0.84, 1.16 | 10.12 | 120 |

Table 1: A summary of the properties of the systems presented, including variety of inter-particle force, particle dispersity, particle sizes, spatial density of particles, \(\phi\), and simple area fractions of particles, \(\Phi\). We note, particles are point particles in simulations, C; hence, diameters are not defined in system C.

4.1 Experiments

Using a custom built interfacial stress rheometer (ISR, SI Fig. 1), we simultaneously measure storage and loss moduli and track particle positions in 2D dense suspensions of athermal, repulsive particles. The ISR measures rheology by imposing force on a magnetic needle adsorbed at an interface between oil and water\(^1\). A stationary wall is opposite the needle, so that shear is imposed over a distance visible by a microscope. The displacement of the rod is measured precisely with the microscope. With displacement (strain) and imposed force (stress), the storage and loss moduli are calculated\(^2,3\). Additionally, the microscope is used to image the particles (\(\sim\)40,000, from wall to needle) adsorbed at the interface. The particles include charges on their surfaces, so they exert dipole-dipole repulsive forces on each other\(^4–6\). At the particle densities in these experiments, these forces result in particle jamming, which we define as full kinematic restraint on each particle by its neighbors. In all data reported here the systems are in a sinusoidal, steady state. In the experiments, steady state occurs after five shear cycles. Twenty-five steady state cycles are used for calculations. For more information about these experiments and the calculations of \(D_{\text{min}}^2\) see Refs.[7–9].

An accessible quantity in our experiments is the two-body approximation of excess entropy, the difference between the system’s entropy and the entropy of an ideal gas in an equivalent state (\(s_2 \sim s_{\text{sys.}} - s_{\text{I.G.}}\)). Conveniently, this quantity is calculated from the radial distribution function, which is available in a wide range of experiments\(^10\). The previously derived\(^11\) formula for excess entropy is:

\[
s_2 = -\pi \rho \int_0^\infty \{g(r) \ln[g(r)] - [g(r) - 1]\} r dr
\]

where \(\rho\) is the particle number density. We implement equation 4 for each image in our experiments individually to collectively construct an entropy time signal, \(s_2(t)\). For specifics of our excess entropy calculations, see Ref. [12].

The network force, \(F^*\) introduced in the paper is calculated based on inter-particle forces within the average neighborhood of particles. To make this measurement we estimate the average number of nearest neighbors around a particle as:

\[
Z(R_c) = 2\pi \rho \int_0^{R_c} g(r)r dr
\]

where \(R_c\) values are shown as the horizontal axis in Fig. 1c. We estimate experimental inter-particle forces based on potentials measured in experiments and molecular dynamics simulations reported in Ref. [6]. An account of our estimate is included in the Supplemental Materials.

4.2 Simulations

The data points for samples C were obtained using LAMMPS\(^13\). At each strain amplitude, 10 two-dimensional ensembles of 10,000 bi-disperse Lennard-Jones particles\(^14,15\) were subjected to sinusoidal shear under periodic boundary conditions at constant confining pressure. The period of shearing was 100\(\times\) that of the LJ time-scale of the particles. Prior to shearing, the samples were dynamically equilibrated at 1% of the glass-transition temperature\(^15\). During strain-controlled shearing LAMMPS’ Nosé-Hoover thermostat was used to maintain the samples at approximately 1% of the glass-transition temperature. After 40 cycles of shearing, the shear stress was output for another 40 cycles
for later use in the calculations of the dynamic moduli. We find that similar calculations at 9% of the glass-transition temperature begin to violate our assumption of negligible thermal energy.

For simulation samples D and E, we use HOOMD-blue\textsuperscript{16,17} to impose cyclic strain on 10 particle configurations for each of six strain amplitudes (1, 2, 3, 4, 5, 6%) at constant confining pressure. Ensembles are composed of jammed states of 50:50 bidisperse mixtures of 10,000 Hertzian particles. Ensembles are initialized from a randomly uniform probability distribution at a packing fraction below jamming, and subsequently quenched under FIRE minimization\textsuperscript{18} whilst increasing the packing fraction until the desired pressure is reached. We then run a triangle wave shear protocol, imposing a small strain step of $10^{-4}$% and minimizing under FIRE after each step, until a total of 40 cycles have been completed. We calculate dynamic moduli based on the dominant frequencies of the resulting triangle waves.

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