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Macrosopic yielding in jammed solids is accompanied by a non-equilibrium first-order transition in particle trajectories

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We use computer simulations to analyse the yielding transition during large-amplitude oscillatory shear of a simple model for soft jammed solids. Simultaneous analysis of global mechanical response and particle-scale motion demonstrates that macroscopic yielding, revealed by a smooth crossover in mechanical properties, is accompanied by a sudden change in the particle dynamics, which evolves from non-diffusive motion to irreversible diffusion as the amplitude of the shear is increased. We provide numerical evidence that this sharp change corresponds to a non-equilibrium first-order dynamic phase transition, thus establishing the existence of a well-defined microscopic dynamic signature of the yielding transition in amorphous materials in oscillatory shear.

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I. INTRODUCTION

A major effort in soft condensed matter physics concerns the design of materials with well-controlled mechanical properties [1]. Rheology thus represents a central probe and oscillatory shear measurements at finite frequency $\omega$ are among the most commonly performed mechanical tests [2]. In this approach, a harmonic deformation is applied and the stress response measured, or vice-versa. In the linear regime, the complex shear modulus $G^*(\omega) = G'(\omega) + iG''(\omega)$ provides information about the nature and strength of the material at a given frequency [3], while microscopic relaxation processes can be probed by varying the frequency. At larger amplitude, non-linear mechanical properties are accessed.

This approach is well-suited for amorphous materials, which often display non-trivial response spectra in the linear regime where they behave as soft elastic solids, but flow as the amplitude of the forcing is increased beyond a “yielding” limit [4–6]. Whereas the storage modulus $G'(\omega)$ dominates the elastic response at small deformation amplitude, irreversible plastic deformations occur post-yielding where the loss modulus $G''(\omega)$ instead dominates. Oscillatory shear experiments have been performed in a wide range of soft condensed matter systems across yielding, such as granular particles [7–11], emulsions [12–15], colloidal suspensions [16–18] and gels [19], as well as in computer simulations [20–24].

In experiments, the change from elastic to plastic response in macroscopic mechanical properties is often described as a “yielding transition”, even though yielding appears as a smooth crossover whose location cannot be unambiguously defined [25]. Interestingly, recent experiments have provided evidence that this macroscopic crossover corresponds to a qualitative change in particle trajectories [9, 11, 12, 18, 26–28]. As expected physically, particles are essentially arrested in the undeformed solid, but can diffuse due to irreversible plastic rearrangements occurring at larger amplitude. There is, however, no consensus about the nature of this crossover, which has been described either as a smooth change [10], as a relatively sharp crossover [12], or as a continuous non-equilibrium phase transition [18]. The latter conclusion builds a qualitative analogy with the continuous irreversibility transition observed in low-density suspensions [7, 8, 29], which has been actively studied in computer simulations [23, 24], and attempts to borrow concepts from the field of non-equilibrium phase transitions [30–32]. In addition recent experiments argue that yielding corresponds to a change in the microstructure of the system [27], by opposition to the dynamic properties discussed here. A clear connection between these microscopic changes and the macroscopic rheology is lacking.

Here, we use a model of a jammed material composed of non-Brownian repulsive spheres to investigate the nature of the yielding transition at the particle-scale level, in the simple situation where thermal fluctuations and hydrodynamic forces play no role. We reproduce standard mechanical signatures of macroscopic yielding under oscillatory shear and obtain two key results regarding particle trajectories. First, we show that the onset of particle diffusion in steady state is extremely sharp and occurs at a well-defined shear amplitude, which unambiguously locates the yielding transition. Second, we find that particle diffusivity emerges discontinuously at yielding, thus demonstrating that yielding corresponds to a non-equilibrium first-order phase transition. These findings differ qualitatively from earlier suggestions of a continuous irreversibility transition [18, 24], but seem to agree very well with recent experimental findings [10, 12]. We also show that this transition is dynamic in nature, and is not accompanied by discontinuous structural changes.

II. MODEL AND NUMERICAL TECHNIQUES

We consider soft repulsive non-Brownian particles in a simple shear flow geometry. We perform standard over-
damped Langevin dynamics simulations of a well-known model of harmonic particles in three dimensions [33, 34], using an equimolar binary mixture of small and large particles with diameter ratio 1.4. The equations of motion read

\[ \dot{\mathbf{r}}_{ij} = \frac{1}{m_{ij}} \left( \frac{\partial U}{\partial \mathbf{r}_{ij}} - \gamma(t) \mathbf{v}_i(t) \right), \]

where \( \gamma \) is a friction coefficient, \( \gamma(t) = (x_i + y_i + z_i) \) is the shear rate. For particles \( i \) and \( j \) having diameters \( a_i \) and \( a_j \), the pair potential reads \( U(r_{ij}) = \frac{1}{2} (1 - r_{ij}/a_i)^2 \Theta(a_i - r_{ij}) \), where \( \Theta \) is an energy scale, \( a_{ij} = (a_i + a_j)/2 \), \( a \) denotes the diameter of small particles, and \( \Theta(a) \) is the Heaviside function. The unit length is \( a \), the unit time \( \tau_0 = a^2 \xi_0 / \epsilon \), the unit energy \( \epsilon \), and so the unit stress is \( \epsilon / a^3 \).

We apply a harmonic deformation using Lees-Edwards periodic boundary conditions [34], the strain evolving as \( \gamma(t) = \gamma_0 [1 - \cos(\omega t)] \), where \( \gamma_0 \) is the amplitude of the imposed shear strain and \( \omega = 2\pi / T \) is the frequency of the oscillation. The period \( T \) is chosen very large to be close to a quasi-static protocol; we use \( T = 10^4 \tau_0 \). We have checked that our results do not qualitatively depend on this choice. We work at constant packing fraction \( \varphi = 0.80 \), much above the jamming density \( \varphi_J \approx 0.647 \) [35]. We checked that our results are representative of the entire jammed phase, \( \varphi > \varphi_J \), but yielding could be more complicated in the limit \( \varphi \to \varphi_J \) where the system looses rigidity [12]. The different regime \( \varphi < \varphi_J \), where a yield stress does not exist, was analysed before [29]. We solve Eq. (1) with a modified Euler algorithm [34], using a discretization timestep \( \Delta t = 0.1 \tau_0 \). Numerical stability and accuracy were carefully checked.

To investigate finite-size effects, we perform simulations with four different sizes \( N = 300, 1000, 3000, \) and \( 10000 \), where \( N \) is the total number of particles. All simulations start from fully random configurations. We analyse both the transient regime after shear is started, and steady-state measurements. To improve the statistics, we perform at least 4 independent runs for each pair \((\gamma_0, N)\).

At the macroscopic level, our main observable is the time-dependent response of the \( y \)-component of the shear stress, defined by the usual Irvnig-Kirkwood formula [34]: \( \sigma(t) = -\frac{1}{2} \sum y_{ij} F_{ij}^y \), where \( V \) is the volume and \( F_{ij}^y \) represents the \( y \)-component of the force \( F_{ij} \). The kinetic part of the stress is fully negligible in the present situation of low-frequency oscillatory shear. To analyse the rheological response in steady state, we fit the time series of the shear stress to a sinusoidal form,

\[ \sigma(t) = -\sigma_0 \cos(\omega t + \delta), \]  \hspace{1cm} (2)

where \( \sigma_0 \) is the amplitude of the first harmonics at frequency \( \omega \), and \( \delta \) is the phase difference between strain and stress. In practice, \( \sigma_0 \) and \( \delta \) are obtained by fitting Eq. (2) to steady state data lasting about 1000\( \tau_0 \). Alternatively, we can transform the two parameters \((\sigma_0, \delta)\) into the more conventional quantities \( G'(\omega) \) and \( G''(\omega) \) using

\[ G'(\omega) = \frac{\sigma_0}{\gamma_0}, \quad G''(\omega) = \frac{\sigma_0 \delta}{\gamma_0^2}. \] \hspace{1cm} (3)

III. SMOOTH CROSSOVER IN MACROSCOPIC RHEOLOGY

In Fig. 1(a), we show the evolution with the strain amplitude \( \gamma_0 \) of the storage and loss moduli at fixed frequency \( \omega \) measured in steady state. At very low \( \gamma_0 \), \( G' \) dominates the response, \( G'/G'' \approx 10 \), indicating that the system responds in the linear regime as a soft elastic
solid. As $\gamma_0$ is increased, the moduli first evolve slowly for $\gamma_0 < 0.1$, where little plastic rearrangements are produced. As $\gamma_0$ increases further, we observe a crossing of $G'$ and $G''$ at $\gamma_x \approx 0.15$ (dashed lines), so that dissipation dominates $\gamma_0 > \gamma_x$. These mechanical properties reproduce well-known behaviour [1, 2] and validate our numerical approach. We notice further that they display virtually no finite-size effects. In this representation, $\gamma_x \approx 0.15$ appears as the most relevant strain scale to characterize yielding, although a smooth crossover can be qualitatively detected near $\gamma_0 \approx 0.1$, where the $\gamma_0$-dependence of the moduli becomes somewhat steeper.

In Fig. 1(b) we plot this rheological evolution in the alternative representation offered by $(\alpha_0, \delta)$. At small $\gamma_0$, the phase $\delta$ is very small while the stress amplitude increases linearly, $\sigma_0 \approx G'\gamma_0$, as expected for reversible elastic deformations in a solid. Near $\gamma_0 \approx 0.1$ two changes are observed. First, $\sigma_0$ ceases to be linear and displays an overshoot when $\gamma_0$ is increased, signaling that plastic events take place. We define the onset of plastic events, $\gamma_0$, as the location of the stress overshoot (dashed lines in Fig. 1(b)). Our interpretation for $\gamma_0$ is reinforced by the evolution of the phase $\delta$ in Fig. 1(b), which grows steadily above $\gamma_0$, indicating the onset of dissipation. Note that the crossing of $G'$ and $G''$ at $\gamma_x$ has no obvious relevance in this representation where it simply corresponds, by definition, to the strain scale where $\delta = \pi/4$. In Fig. 1(c) we confirm that $\gamma_0$ and $\gamma_x$ display virtually no system size dependence, but that they differ quantitatively.

Whereas $\gamma_x$ is frequently quoted as “the” yielding point in the literature, a stress overshoot also serves to identify yielding in shear-start experiments [25]. A stress overshoot is reported in some oscillatory shear experiments [19], but is absent in others [12, 13]. A possible explanation is that experiments are typically performed at somewhat larger frequencies, where additional contributions to the shear stress (lubrication forces, hydrodynamic effects) might hide this behaviour. In addition, we have confirmed that the overshoot disappears and is replaced by a monotonic increase seen in experiments [13] when we use a substantially larger frequency, typically $\omega > 10^{-3}$. We emphasize that the presence of the dynamic transition discussed below is independent of the existence of the stress overshoot reported in Fig. 1(b).

IV. SHARP TRANSITION IN MICROSCOPIC DYNAMICS

We now turn to the evolution of single particle dynamics. A first natural dynamic observable is the averaged particle displacement after one deformation cycle [12, 24],

$$\Delta r(t, T) = \frac{1}{N} \sum_j |\vec{r}_j(t + T) - \vec{r}_j(t)|,$$  \hspace{1cm} (4)

where $t$ is the time since shear is applied. In Fig. 2(a), we show how $\Delta r(t, T)$ evolves in the transient regime for various amplitudes of the applied deformation. For small $\gamma_0$, particle displacements decay rapidly to zero. In the elastic solid at small amplitude, there are rare rearrangements taking place before the system settles near a stable energy minimum where particles have nearly periodic motion (or quasi-periodic motion with a period that is a multiple of $T$), as reported before [23, 36]). As $\gamma_0$ is increased, it takes more and more time for $\Delta r(t, T)$
to eventually vanish. When \( \gamma_0 \) is larger than \( \gamma \sim 0.095 \), the average particle displacement never vanishes in the explored time window, but instead fluctuates around a well-defined finite value, which increases with \( \gamma_0 \). This regime corresponds to irreversible, non-periodic particle trajectories. In Fig. 2(b), we plot the time averaged displacement for one cycle \( \langle \Delta r(T) \rangle \) in steady states for various strain amplitudes and system sizes. From \( \langle \Delta r(T) \rangle \), a very clear discontinuous jump is observed between the irreversible and reversible states near \( \gamma_c \). Very close to the transition, the displacements exhibit fluctuations around a well-defined value both above and below \( \gamma_c \). Whereas these fluctuations are infinitely long-lived above \( \gamma_c \), they are only metastable below \( \gamma_c \) before the system finds a reversible state where the displacements become very small. We report the value of \( \langle \Delta r(T) \rangle \) for \( N = 10^4 \) for these metastable states in Fig. 2(b). Overall, these fluctuations appear qualitatively distinct from the algebraic decay observed close to continuous irreversibility transitions [8], and are much closer to the phenomenology observed near discontinuous, first-order phase transitions where metastable phases can be observed over long times. In particular, it appears impossible to describe the decrease of \( \langle \Delta r(T) \rangle \) with a continuous vanishing at the critical value of \( \gamma_{\text{dyn}} \).

Stronger evidence of such a phase transition is obtained from the evolution of the average lifetime of the metastable irreversible phase \( \tau_d \), as depicted in Fig. 2(c) for various system sizes. These data confirm that \( \tau_d \) increases rapidly close to \( \gamma_0 \approx 0.1 \). For larger \( \gamma_0 \), trajectories remain irreversible. By contrast to the rheology, a clear system size dependence is observed, larger systems take more time to settle in a global energy minimum. A diverging lifetime is typically observed close to non-equilibrium phase transitions [8, 30, 31], and was reported before [23, 24]. Such divergence is expected for both a continuous or discontinuous transition (see [37, 38] for recent examples). Solid lines in Fig. 2(c) represent an empirical fit, \( \tau_d \sim (\gamma_{\text{dyn}} - \gamma_0)^{-\alpha} \), with \( \alpha \approx 2.1 - 3.0 \) [39], suggestive of a divergence of \( \tau_d \) when approaching the dynamic transition at \( \gamma_{\text{dyn}} \). We notice that finite size effects can be felt even when the system is not very close to the critical value, such that we cannot observe the expected saturation of \( \tau_d \) to a finite value as \( N \to \infty \). The system size dependence of \( \gamma_{\text{dyn}} \) reported in Fig. 1(a) is very modest and seems to extrapolate to a finite value, \( \gamma_{\text{dyn}} \approx 0.095 \), as \( N \to \infty \).

We now characterize the steady-state irreversible dynamics at large \( \gamma_0 \) using the mean-squared displacement,

\[
\langle \Delta r(t)^2 \rangle = \frac{1}{N} \sum_{j=1}^{N} |r_j(t) - r_j(0)|^2,
\]

where the brackets indicate a time average. The results are displayed in Fig. 3(a) for \( N = 10^4 \), for time delays commensurate with the period. The dynamics is diffusive at long times, \( \langle \Delta r(t)^2 \rangle \sim 6 D t \), where \( D \) is the diffusion constant. We represent \( D \) (in units of \( a^2 / T \)) for various \( \gamma_0 \) and \( N \) in Fig. 3(b). As expected, \( D = 0 \) below a critical value \( \gamma_c \), corresponding to the phase characterized by quasi-periodic particle trajectories, and it increases with \( \gamma_0 \) above \( \gamma_c \). Both \( D \) and \( \Delta r(t, T) \) in Eq. (4) could serve as order parameters for the transition. By measuring \( \gamma_c \) for various system sizes, we observe a modest change with system size, see inset of Fig. 3(b), suggestive of a finite limit \( \gamma_c \approx 0.0885 \) for \( N \to \infty \). The functional form of our extrapolation should be confirmed by additional larger scale simulations.

A striking finding in Fig. 3(b) is the finite amplitude of the diffusion constant at the transition. Near continuous irreversibility transitions, \( D \) decreases by several orders of magnitude and scales algebraically as \( \gamma_c \) is approached from above [7, 40]. We observe instead a modest decrease of \( D \), followed by a sudden jump to zero, which is robust against finite-size effects. In particular we find that diffusive behaviour also persists for a finite amount of time below in the reversible phase, as also described above for the one-cycle particle displacement \( \Delta r(T) \). It is how-
ever more difficult to measure $D$ in this region, because a careful determination of $D$ requires taking the long-time limit, which is not possible by construction in the metastable region. We conclude therefore that the discontinuous behaviour of $D$ in Fig. 3(b) appears less convincing than the one of $\Delta(r,T)$ shown in Fig. 2(b), but the overall phenomenology reported in this work appears inconsistent with a continuous transition.

V. NO CHANGE IN MICROSCOPIC STRUCTURE

It was recently argued that the yielding transition in oscillatory shear can be detected through the static structure of the system [27]. Such a behaviour would differ qualitatively from our conclusion that yielding is revealed through the dynamic evolution of the system. Our analysis of the pair correlation function across the yielding transition did not reveal any change in the static properties of the system in the two phases, which seems to contradict the results of Ref. [27]. To reinforce this conclusion, we have measured the exact same quantity that was detected experimentally. In detail, we resolve the radial dependence of the static structure factor $S(q) = \frac{1}{N} \langle \rho(q) \rho(-q) \rangle$ in the $(x,z)$ plane [27], where $\rho(q)$ is the Fourier component of the density at wavevector $q$. Thus we obtain the $\alpha$-dependence from $\tan \alpha = (q_x/q_z)$, and follow the $\alpha$-dependence of the static structure, as proposed in [27].

To obtain statistically reliable data close to the dynamic transition, we perform an extensive time average over 100 well separated times for the diffusive phase at $\gamma_0 = 0.12$. For the non-diffusive phase at $\gamma_0 = 0.10$, time averages are not useful and we obtain instead 100 independent configurations starting from independent initial conditions. Errorbars are defined from the resulting sample-to-sample fluctuations. Because the stress is close to a sinusoidal form, we measure the structure either when the stress is zero ('undeformed' states) and when it is maximal $\sigma(t) = \pm \sigma_0$ ('deformed' states). Thus, we obtain 4 measures of the structure at two shear amplitudes, for both deformed and undeformed states.

In Figure 4(a), we show the $q$-dependence of the averaged structure factor $S(q) = |q|$ for these 4 cases and for $N = 1000$. We observe that neither the particle reversibility nor the deformation seem to affect much the structure factor. We now resolve the angular dependence of $S(q,\alpha)$ using wavevectors in the $(x,z)$ plane having an amplitude close to the first peak in the range $q_0 = 5.9 - 7.0$. In Fig. 4(b), we show that the $\alpha$-dependence of the structure factor for the 4 situations defined above is essentially inexistente. Importantly, we do not observe any difference for reversible and irreversible regimes across yielding. In particular, we do not observe the oscillations that were detected in the experiments for the arrested phase. Furthermore, we also checked that average values of other static quantities (such as the energy density and pair correlation functions) are similarly insensitive to the underlying dynamic transition. These conclusions contrast with the results in Ref. [27], which are perhaps due to the larger shear rates employed in the experiment. Another major contradiction with that work is our finding that yielding does not correspond to the crossing point of $G'$ and $G''$. Our results show that yielding is best interpreted as a loss of reversibility in the particle trajectories, which is a purely dynamical concept.

VI. CONCLUSION

Together, our results suggest that the yielding transition of jammed solids under large-amplitude oscillatory shear is accompanied by a first-order non-equilibrium phase transition, rather than a continuous one. It marks
the abrupt emergence of irreversible non-affine particle motion. The characteristic strain amplitudes obtained from rheology ($\gamma_x, \gamma_pl$) and from microscopic dynamics ($\gamma_{dyne}, \gamma_c$) are compiled in Fig. 1(c). To a good approximation, we find $\gamma_pl \approx \gamma_{dyne} \approx \gamma_c$, whereas $\gamma_x$ is significantly larger, corresponding to a large amount of dissipated energy. The dynamic phase transition revealed by the discontinuous evolution of single-particle dynamics produces a smooth crossover in mechanical properties at a critical strain amplitude that appears unrelated to the crossing of $G'(\omega)$ and $G''(\omega)$. Our conclusions contrast with earlier claims of a continuous transition [18, 24], but appear in very good agreement with observations in a sheared emulsion [12]. We hope our study will trigger further work in a broader variety of numerical and experimental systems to fully establish its generality.

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