Pre-Yield Non-Affine Fluctuations and A Hidden Critical Point in Strained Crystals

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A crystalline solid exhibits thermally induced localised non-affine droplets in the absence of external stress. Here we show that upon an imposed shear, the size of these droplets grow until they percolate at a critical strain, well below the value at which the solid begins to yield. This critical point does not manifest in most thermodynamic or mechanical properties, but is hidden and reveals itself in the onset of inhomogeneities in elastic moduli, marked changes in the appearance and local properties of non-affine droplets and a sudden enhancement in defect pair concentration. Slow relaxation of stress and an-elasticity appear as observable dynamical consequences of this hidden criticality. Our results may be directly verified in colloidal crystals with video microscopy techniques but are expected to have more general validity.

Mechanical properties of solids¹², especially mechanisms of yielding in response to external stress, continue to engage the attention of materials scientists and engineers⁴. Pre-yield phenomena such as anelasticity, occurring at non-zero temperatures and below the yield stress, in both crystals⁴ and metallic glasses⁵,⁶ is somewhat less understood, however. Within this anelastic regime, a crystalline solid undergoes recoverable strain but with a long relaxation time and is accompanied by conspicuous production and reorganisation of lattice defects⁷,⁸ marked by the onset of significant departure from instantaneous and linear, "Hooke's law", elasticity. Mechanical response becomes heterogeneous: the regions with defects behaving differently from the rest of the solid. In amorphous solids too, anelasticity is marked by the appearance of similar heterogeneous mechanical response confined to localised clusters of particles undergoing large non-affine deformation or non affine droplets⁹–¹¹.

In this paper, we focus on defect nucleation and associated pre-yield phenomena in two (2D) and three-dimensional (3D), initially homogeneous, crystals of interacting particles. We provide strong evidence associating the proliferation of crystalline defects during deformation at non-zero temperatures, with the critical behaviour of thermally generated non-affine droplets. These non-affine displacements exist in crystalline solids even in the absence of an applied strain and have a purely thermal origin¹². Upon shearing the crystalline solid, these clusters grow and eventually percolate at a hidden mechanical critical point; concomitantly, defects suddenly proliferate. Significantly, this happens well before the yield point of the crystalline solid. Apart from its intrinsic novelty, this viewpoint sheds insight onto a broader set of issues. First, we find that dislocation nucleation is stochastic and seeded from regions with large non-affine displacements. Second, we have identified the localised non-affine regions with droplet excitations from nearby metastable liquid/glass¹³. Thus, our finding that the percolation of non-affine droplets coincides with the disappearance of the metastable liquid-glass spinodal, brings out the significance of

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metastable configurations in understanding mechanical properties of solids. Last, this perspective might provide a language bridging ideas concerning the mechanical response of crystals and amorphous solids.

The existence of a unique reference in crystals allows us to identify localized non-affine regions at non-zero temperatures, even at zero applied strain. In Ref. 13 we studied the statistics of shape and size of non-affine clusters and their local thermodynamic properties using molecular dynamics (MD) simulations of an unstrained, two-dimensional Lennard-Jones (2D-LJ) solid\textsuperscript{14,15}. We summarise the main results below for completeness.

Even without external strain, thermal fluctuations ensure that local sub-volumes within the crystaline solid undergo random distortions about their equilibrium configurations. For every configuration, particles undergoing large non-affine displacements are identified using the parameter $\chi_i$, viz. the least square error incurred in trying to fit an instantaneous local volume, $\Omega_i$, to an affine distortion of the same volume in the, reference, undistorted lattice\textsuperscript{11}. We choose $\Omega$ to correspond roughly to the bare elastic correlation length or a couple of near neighbour shells. A cutoff criterion $\chi_i > \chi_{\text{cut}}$ then eliminates trivial harmonic distortions of the lattice (see Methods and supplementary information SI). These particles are observed to cluster together into droplets.

The droplets are characterized by a distribution of the local density $\rho_i$ and excess pressures, $\Delta p_i \equiv p_i - p$. Here $p$ is the mean pressure of the surrounding solid of density $\rho$ and $p_i$ is computed by averaging the virial over the $n_i$ particles of the cluster. For any temperature $T$ the mean scaled excess pressure $\bar{p} = \Delta p_i / T$ vs $\rho_i$ curve is non-monotonic and resembles a van der Waals loop seen in usual liquid-gas transitions. The two stable branches with $\partial \bar{p} / \partial \rho_i > 0$ are connected by an “unstable” branch where $\partial \bar{p} / \partial \rho_i < 0$. In the unstrained solid the stable branches were associated with inflated, compact clusters or deflated, string-like clusters depending on whether $\bar{p}$ is negative or positive respectively\textsuperscript{16,17}; the unstable branch contains only ramified “branched-polymer” clusters\textsuperscript{18}. By studying the local equation of state and density-correlations, one may associate compact and string-like clusters with droplet fluctuations from nearby liquid and glass-like metastable free-energy minima respectively\textsuperscript{13}. As $T$ is increased, the van der Waals loop as well as the distinction between compact and string-like droplets vanish above a metastable critical point. This behaviour is quite general and exists in other model systems admitting non-crystalline phases, (see discussion in SI).

Here, we extend this earlier analysis\textsuperscript{13} to the mechanical response of LJ solids under external strain in two and three dimensions over a range of $\rho$ and for a few $T$. In what follows, the bulk of our quantitatively detailed results, unless otherwise stated, are explicitly demonstrated for the 2D-LJ solid for which computations, as well as experimental verification using real-time video microscopy\textsuperscript{19}, are relatively cheap. Key results for the 3D-LJ case\textsuperscript{20} are also presented showing that our main conclusions carry over to higher dimensions.

**Results**

**Percolation of clusters.** As the crystal is subject to a quasi-static (pure) shear strain $\varepsilon = \varepsilon_{xx} - \varepsilon_{yy}$ (Fig. 1a–c) (see Methods), localized non-affine deformations grow and increase in number. At a critical value of the strain $\varepsilon^*$ (e.g. $\approx 0.05$ for $\rho = 0.99$, $T = 0.4$), regions containing non-affine particles begin to percolate. Further increase of the strain beyond $\varepsilon^*$, eventually leads to yielding of the solid (at $\varepsilon \approx 0.1$ for same $\rho$, $T$).
To ascertain the nature of this percolation transition, we plot the ratio of the number of particles in the largest cluster to the total number of non-affine particles, $f_\phi$, in the 2D-LJ solid as a function of the fraction of non-affine particles $\phi$ ($\rho, \varepsilon$) in Fig. 2a. The collapse of $f_\phi$ onto a single curve for all $\rho$ and $\varepsilon$, is a strong indication of a true critical percolation transition at a value of $\phi^* \approx 0.4$, close to the known value of site percolation in 2D (0.34). We plot the probability distribution $P(n_c)$ of obtaining a non-affine cluster of size $n_c$ for increasing values of $\varepsilon$ in Fig. 2b. As one approaches the percolation transition, $P(n_c) \sim \tau$ with an exponent $\tau \approx 1.67$.

We now ask whether there is a signature of this geometrical critical point in the thermodynamics and whether these non-affine clusters admit a thermodynamic interpretation. The answer to this is surprisingly subtle.

**Local thermodynamics.** Bulk thermodynamic quantities such as equation of state etc. show no abrupt change across the transition. We, therefore, focus on measuring local thermodynamic variables within the non-affine clusters and monitoring them across the percolation transition. We characterise droplets using the local quantities $\bar{\rho}$ and $\rho_c$ (see also Methods). The local temperatures, on the other hand, are close to the fixed temperature of the solid and are irrelevant to the subsequent analysis. Unlike Ref. 13, instead of temperature, here strain, $\varepsilon$, drives the percolation transition.

We anticipate that systems with various $\varepsilon$ and $\rho$ behave similarly for equal $\phi$. We test this as follows. From configurations with fraction of non-affine particles between $\phi$ and $\phi + \phi$, where $\Delta \phi$ is a suitable bin-size, we obtain the mean $\bar{\rho}$ and $\rho_c$ corresponding to droplets with fixed number of particles $10 < n_c < 100$; collecting together data from all solids with $0.91 \leq \rho \leq 0.99$ and $\varepsilon$ below the yield point. This data is plotted in Fig. 3a for droplets of size $n_c = 15$ for each value of $\phi$. Similar plots are obtained for droplets with other values of $n_c$ too.

Over a large range of $\phi$ values, the plots are non-monotonic, showing prominent van der Waals loops with two stable branches. We show below that these correspond to compact and string-like droplets. As $\phi$ increases, the loops tend to vanish. Within the accuracy of our computations, the value of $\phi$ above which the loops vanish is the same as $\phi^*$, i.e. the non-affine fraction at which the clusters percolate.

To make this identification quantitative, we obtain the values of $\rho_c$ for all $\phi$ at the spinodal $d\bar{\rho}/d\rho_c = 0$ and plot them in Fig. 3b for $n_c = 10, 15$ and 20. When extrapolated, the spinodal lines intersect at a metastable, droplet critical point. This metastable critical point and the droplet shape transition survives for even higher values of $n_c$, though for very large $n_c$ the statistics becomes sparse.

How does the extent of the van der Waals loop in $\bar{\rho}$ vs. $\rho_c$ depend on the size of the cluster $n_c$? In Fig. 4a we show a comparison of the van der Waals loops for various values of $n_c$ in the $\varepsilon \rightarrow 0$ limit. An integration of $\bar{\rho}$ vs. $\rho_c$, which gives a measure of the free energy $\mathcal{F}$, expended by the solid to produce these fluctuations has a convex up region, showing that an interpretation in terms of a metastable first order transition is not inconsistent. A metastable first order transition implies, at least for compact
Figure 3. a Plot of $\bar{p}$ vs. $\rho_c$ for various $\phi$ (key on right) and $n_c = 15$ showing van der Waals loops. Loops for different $\phi$ have been shifted in $\bar{p}$-axis for clarity. Note that the loop disappears for $\phi > \phi^*$. b Plot of the spinodal obtained for $n_c = 10, 15$ and 20 in the $\phi - \rho_c$ plane clearly indicating a metastable critical point coinciding with $\phi^*$.

Figure 4. a Plot showing $\Delta p_c$ as a function of the local density $\rho_c$ for “non-affine” droplets in the strain free LJ solid at $T = 0.4$. The different colors show the van der Waals loop for various sizes of clusters $10 < n_c < 30$. The data becomes sparse for much higher values of $n_c$, leading to worse statistics. The van der Waals loops however show stabilisation with larger $n_c$. We have made this quantitative by obtaining an effective barrier height $\Delta F_c$ for this transition by integrating the pressure difference $\Delta p_c$ along $\rho_c$ for each of these curves. The result is plotted in b and shows that $\Delta F_c$ increases with $n_c$ as expected of a first order transition. The solid line is a linear fit.
droplets, that the surface free energy barriers should scale as $\Delta F_c \sim n_c^{-\nu}$, where $n_c$ is a typical linear size associated with the clusters and $d (=2)$ is the dimensionality$^{22}$. The actual form of the scaling may be more complicated because of the complex shape of the clusters. Nevertheless, in Fig. 4b we show a plot of the barrier height $\Delta F_c$, obtained by integrating Fig. 4a, as a function of the size of the fluctuation. A monotonically increasing curve indicates a positive surface energy of the droplets at $\phi = 0$. At the metastable critical point we expect this surface energy to vanish, although large fluctuations and strong finite size effects makes an explicit evaluation of this quantity near $\phi = \phi^*$ computationally impractical.

The shape transition. We have shown that the van der Waals loop in $p$ vs $\rho_c$ vanishes as $\phi > \phi^*$ where the non-affine droplets percolate. The van der Waals loop corresponds to a metastable transition between droplets which come in two stable shapes, either compact or string-like. We show now that this distinction between these shapes also vanishes for $\phi > \phi^*$. Above this transition, string-like droplets connect the adjacent compact "nodes" in a single giant percolating network. This network has an overall, branched polymer shape so that isolated compact and string-like droplets cease to exist beyond $\phi > \phi^*$ (see Fig. 1c).

For the unstrained solid$^{13}$ the $\bar{p} < 0$ and $\bar{p} > 0$ branches of the van der Waals loop associated with inflated, compact droplets and deflated, string-like droplets$^{16,17}$ respectively are characterised by the crossover of the mean radius of gyration $R_g(n_c, \Delta p_c, T) = n_c^{\nu} F(\bar{p} n_c^{1/2})$, where $F(x)$ is a crossover function. The value of $\nu$ is 0.64 for branched polymers and the $-ve$ and $+ve \bar{p}$ branches crossover to $\nu = 0.5$ and 1 respectively. In Fig. 5a we recover this crossover of the droplet shapes for $\phi \leq \phi^*$ for the two corresponding branches of the van der Waals loop as in the unstrained solid. As $\phi$ increases, however, the giant percolating cluster dominates the droplet configuration space and it is impossible to obtain compact or string-like droplets; all droplets now show branched polymer behaviour (Fig. 5b) with $R_g \sim n_c^{0.64}$ typical of a system above a critical point. This disappearance of the transition in shapes coincides with the disappearance of the van der Waals loop in $\bar{p} - \rho_c$ plane.

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**Figure 5.** The radius of gyration $R_g$ as a function of the size $n_c$ of the droplets where we have factored out the $\sim n_c^{\nu}$, with $\nu = 0.64$ (branched polymer) dependence. Below the percolation transition, a, the droplets for the $\bar{p} > 0$ and $\bar{p} < 0$ branches behave differently assuming string-like ($\nu = 1$) and compact ($\nu = 1/2$) configurations respectively. Above percolation b, the morphology of droplets changes drastically and all droplets become branched polymers.
Nucleation of defects. Non-affine droplets are associated with coordination number changing deformations. To show this, we track the local concentration of defect pairs by counting the number of nearest neighbours of particles using a local Delaunay triangulation (see Fig. 6a inset). In Fig. 6a, we plot the density of defect pairs $\rho_d$ as a function of the external strain for a LJ solid. While a small number of dislocation pairs (dipoles) always exist within a solid at finite temperatures, there is a sudden increase in $\rho_d$ as soon as $\varepsilon$ crosses a critical value. This increase in defect pair concentration coincides with the critical percolation transition of the non-affine clusters. That this is not a mere numerical coincidence is obvious from the scaling collapse of $\rho_d$ for all $\rho$ and $\varepsilon$ onto a single curve when plotted against the non-affine number fraction $\phi$ in Fig. 6b.

Figure 6. a Plot of the defect concentration $\rho_d$ (particles with 7 Delaunay neighbours) vs. $\varepsilon$ in the 2D-LJ solid at $\rho = 0.99$ showing sharp growth of $\rho_d$ for $\varepsilon > \varepsilon^*$. Inset shows a particle neighbourhood at two different values of $\varepsilon = .039$ (open circles) and .068 (filled gray circles), showing the nucleation of a defect pair (black filled circles). Dashed and the bold lines shows change in topology of the local Delaunay neighbourhood with strain. b Plot of $\rho_d$ vs. $\phi$ for all $\rho$ and $\varepsilon$ showing data collapse onto a single curve with $\rho_d$ increasing sharply at $\phi^*$. Inset shows the same data in semi-log axes to emphasise the small $\phi$ region. Symbols: same as in Fig. 2a. c Particles coloured according to the value of the local non affine parameter $\chi$ for an isolated dislocation-anti-dislocation pair in a triangular crystal. d The same configuration as in c showing non-affine particles with $\chi > \chi_{cut}$ (red filled circles). Majority of non-affine particles are associated with the dislocation dipole (black arrows).
We have obtained identical results also at $T = 0.35$ with data collapse occurring for both $\rho$ and $T$ implying that Fig. 5b represents an universal relation between $\rho_d$ and $\phi$ for all $\rho$, $T$ and $\varepsilon$ for the 2D-LJ solid. A similar phenomenon occurs for the 3D-LJ solid as well. We discuss these results later in this work.

Dislocations are preferentially nucleated within the percolating cluster. Also, each defect pair is dressed with an extended region of non-affine particles contributing to the system-spanning, non-affine cluster. We show this by introducing a dislocation pair deleting a row of 20 atoms in a triangular crystal of $100 \times 100$ LJ atoms at $\rho = 0.97$. After relaxation to obtain a reference configuration, $\chi$ is calculated during a MD run at $T = 0.4$ (Fig. 6c,d). Percolation of non-affine clusters and nucleation of dislocation pairs occur hand-in-hand at the percolation point. This is the main result of the present work.

**Elastic nonlinearity, heterogeneity, anelasticity.** How does this percolation transition affect mechanical properties of the solid? Increased non-affine fluctuations at defect sites is expected to reduce the local elastic modulus $E$. The average elastic modulus therefore involves mechanical properties of the solid. Increased non-affine fluctuations at defect sites is expected to reduce $E$.

During a MD run at $T = 0.4$ (Fig. 6a), we show quasi static stress-strain curves for the LJ solid at $T = 0.4$ for $0.9 < \rho < 0.99$ scaled by $\sigma^*$ and $\varepsilon^*$ the stress and strain respectively at percolation. Notice that for all $\varepsilon \leq \varepsilon^*$, the curves collapse to the trivial, linear, response while for larger strains the response becomes non-linear. The non-linearity $\Delta$ defined as the deviation of the stress from a linear fit to the small strain regime is shown in Fig. 7a. Notice that $\Delta$ changes sign close to $\varepsilon^*$; $\Delta = 0$ for a Gaussian. Here $\sigma$ and $\Sigma$ are the mean and standard deviation of $P(\sigma_b)$.

![Figure 7](image-url)

**Figure 7.** a Scaled stress $\sigma/\sigma^*$ vs. $\varepsilon/\varepsilon^*$ in the 2D-LJ solid at $T = 0.4$. The symbols, which have the same meaning as in Fig. 2a, show values for different $\rho$. Inset shows non-linearity $\Delta$ vs. $\varepsilon$. The vertical dashed line marks $\varepsilon^*$. b Distribution of local stress, $P(\sigma_b)$ obtained by dividing the simulation cells into 1600 equal sub-blocks. At small strains the distribution is Lorentzian. As $\varepsilon$ crosses $\varepsilon^*$, however, spatially correlated dislocation structures develop making sub-blocks. At small strains the distribution is Lorentzian. As $\varepsilon$ crosses $\varepsilon^*$, however, spatially correlated dislocation structures develop making sub-blocks.
at $\varepsilon = 0$. As $\varepsilon$ increases, so does $\rho_d$, making $P(\sigma_B)$ broader. At $\varepsilon > \varepsilon^*$, however, there is a dramatic increase of $\rho_d$ and large numbers of correlated defect pairs are produced. The transformation of $P(\sigma_B)$ can be quantified by calculating the 3rd and 4th order moments Fig. 7b. The average size of the non-affine clusters is expected to remain close to that of $\Omega$ well before the percolation transition and should begin to diverge near the transition quickly reaching the size of the simulation box. Any probe which measures changes of local physical quantities, such as local stress, at the level of $\Omega$ would therefore immediately signal the commencement of percolation. Indeed, at the percolation point the stress distribution rapidly changes from being a sharp Lorentzian to a flat, “sub-Gaussian” form. Position correlation between defects can also be seen directly from particle snapshots (see Fig. 8a-c) as well as the pair distribution function of 7—coordinated particles for solids (see Fig. 8d) as $\phi$ is increased across the transition at fixed $\rho$ and $T$.

Lastly, crystalline solids, as well as bulk metallic glasses, often show complex, time-dependent, stress relaxation behaviour at values of external strains much below the yield point without permanent plastic deformation. To study the an-elastic response of the LJ solid, we quench a few configurations of our LJ solid, which were equilibrated at several values of $\varepsilon$, by rescaling the particle coordinates and the boundary to zero strain. Once initial transients die out, the solid enters a regime of slow relaxation. The nature of this relaxation of the stress, $\sigma(t)$, which is driven by the motion and subsequent annihilation of defect pairs of opposite sign, depends on the number of such defects and therefore on the initial strain. If the initial strain (and consequently, the defect density) is small, this regime is not very prominent. On the other hand, if the initial $\varepsilon$ is large, there are a large number of defects which reorganise over a much longer timescale. In this case, therefore, the second relaxation, as shown in Fig. 9 is complex and shows non-monotonicity. We characterise this relaxation by plotting the time average of the stress $\sigma_a = \tau^{-1} \int_0^\tau dt \sigma(t)$ for various values of the initial strain and observe a sharp change of slope at $\varepsilon^*$ (see Fig. 9, inset). Since defect pairs are formed at the percolation critical point, the origin of non-monotonic relaxation is traced to the percolation transition at $\phi^*$. The anelastic regime, grows with lowering $\rho$ (and increasing $T$) as expected.

**Effect of temperature.** The percolation threshold depends on $\phi$ the fraction of non-affine particles and not separately on temperature, density and strain. The main thesis of our work is that mechanical

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**Figure 8.** a–c Particle snapshots coloured by the value of the local $\chi$ parameter for $\rho = 0.99$ 2DLJ solid at $T = 0.4$ and $\phi = 0.299$ (a), 0.357 (b) and 0.425 (c) showing the increase of spatial correlation of the defects as the solid is strained across the percolation transition at $\phi \approx 0.4$. d The pair distribution function of 7—coordinated particles for the same parameters as in a–c.
Figure 9. Anelastic response of the solid at $\rho = 0.99$. A few strained configurations ($\varepsilon = 0.016, 0.028, 0.036, 0.048, 0.060, 0.064$) are quenched suddenly to zero strain by rescaling the atomic coordinates to study the relaxation of stress $\sigma(t)$ over time $t$. The stress responds by first decreasing rapidly to a negative value whose magnitude increases with the initial $\varepsilon$ and then slowly relaxing to zero from below. For the minimally strained solid, the relaxation is essentially exponential. If the initial strain is around $\varepsilon^*$, the dynamical response shows a second relaxation which is complex and non-monotonic. The inset shows the dependence of the average stress $\sigma_t$ during the relaxation process on the value of the initial $\varepsilon$ before $\sigma(t)$ decreases to 0. While $\sigma_t$ grows linearly for small $\varepsilon$, its growth becomes rapid as $\varepsilon$ crosses $\varepsilon^*$.

properties, such as the defect nucleation threshold, is determined by percolation of non-affine droplets. Therefore, mechanical properties of the solid should depend on external parameters such as temperature, density and strain only through the dependence of $\phi$ on these parameters. We have shown this in Fig. 2a,b for the combination of density and strain at a fixed temperature. It is clear that $\phi$ decreases with increasing density and increases with strain. The percolation threshold at fixed temperature therefore moves to higher strain with increasing density and lower density with increasing strain. If, on the other hand, the temperature is lowered at fixed density one should go to higher strains to achieve percolation. Similarly, the percolation threshold, for fixed strain, should move to lower densities at lower temperatures.

To put to test the validity of our results at other temperatures, we have computed $\rho_d$ for a 2DLJ solid with $\rho = 0.93$ and 0.99 at both $T = 0.35$ and $T = 0.40$. We show that $\phi (T, \rho)$ in Fig. 10a behaves exactly according to our expectations. We also show in Fig. 10b that one can collapse the data for $\rho_d$ as the solid is quasi-statically strained at different temperatures and densities using the non-affine fraction $\phi$. The relation shown in Fig. 6b is therefore, indeed, an universal curve with $\phi$ vs. $\varepsilon$ and the stress-strain curves collapse onto each other when scaled by $\varepsilon^*$ and $\sigma^*$. The cause for nontrivial mechanical response is the nucleation of defects which increase rapidly for $\varepsilon > \varepsilon^*$ where clusters containing non-affine particles percolate. Are our results valid only for the 2D-LJ solid or does this have more general validity? To check this, we have looked at the 3D-LJ solid which has a close packed FCC structure. Since fluctuations generally decrease with increase in the number of dimensions, a naïve argument may suggest that non-affine fluctuations may not have much of a role in determining the mechanical response of the 3D-LJ solid. The situation is however more subtle, since in higher dimensions there are more modes that are non-affine. For example, choosing a neighbourhood which includes only nearest neighbours, the triangular lattice with 6 neighbours, features 4 affine (volume, uniaxial extension, shear and rotation) and 8 non-affine modes$^{12}$, while in three dimensions, the FCC lattice with 12 neighbours has 9 affine and 27 non-affine modes. It is possible that the larger number of available non-affine modes more than compensate for the reduction of fluctuations due to the increased dimensionality.

We simulate the 3D-LJ solid (see Methods) at LJ reduced temperatures of $T = 0.7$ and 0.8 and at $\rho = 1.5$ and 1.2. The fraction of non-affine particles $\phi$ is obtained using a procedure analogous to what we use in 2D (see Fig. 11a-c). The solids are subjected to deviatoric strain $\varepsilon = \varepsilon_{xx} - \varepsilon_{yy} - \varepsilon_{zz}$ and the conjugate stress $\sigma$, $\phi$ and $f_\phi$ obtained as in 2D. As expected, $\phi$ increases with $\varepsilon$ for fixed $\rho$ and $T$ and decreases at fixed $\varepsilon$ as either $\rho$ is decreased or $T$ increased similar to the 2D case; $f_\phi$ vs. $\phi$ is an universal curve with

Extension to three dimensions. We have shown that in the 2D-LJ solid at any density and temperature, defect nucleation happens at a particular value of strain $\varepsilon^*(\rho, T)$ where non-affine clusters percolate. The mechanical response of the solid $\sigma(\varepsilon)$ is trivial below $\varepsilon^*$ and the stress-strain curves collapse onto each other when scaled by $\varepsilon^*$ and $\sigma^*$. The cause for nontrivial mechanical response is the nucleation of defects which increase rapidly for $\varepsilon > \varepsilon^*$ where clusters containing non-affine particles percolate. The percolation threshold at fixed $\rho$ and $T$, $\phi (\varepsilon, \varepsilon^*)$ where clusters containing non-affine particles percolate. Are our results valid only for the 2D-LJ solid or does this have more general validity? To check this, we have looked at the 3D-LJ solid which has a close packed FCC structure. Since fluctuations generally decrease with increase in the number of dimensions, a naïve argument may suggest that non-affine fluctuations may not have much of a role in determining the mechanical response of the 3D-LJ solid. The situation is however more subtle, since in higher dimensions there are more modes that are non-affine. For example, choosing a neighbourhood which includes only nearest neighbours, the triangular lattice with 6 neighbours, features 4 affine (volume, uniaxial extension, shear and rotation) and 8 non-affine modes$^{12}$, while in three dimensions, the FCC lattice with 12 neighbours has 9 affine and 27 non-affine modes. It is possible that the larger number of available non-affine modes more than compensate for the reduction of fluctuations due to the increased dimensionality.

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a percolation transition at $\phi \approx 0.19$, close to the expected value for 3D site percolation\(^{21}\) (see Fig. 12a). Using these results, we obtain $\varepsilon^*$ and $\sigma^*$, the critical strain and conjugate stress at percolation and use this to scale the stress-strain curve shown in Fig. 12b. Again, as in 2D, the stress-strain response becomes non-trivial only beyond the point that non-affine droplets percolate.

Before we end this section, we point out a remarkable aspect of the data shown in Fig. 12. In 3D, identification of the dominant topological defect is non-trivial. In the FCC lattice, lattice dislocations, frequently decompose into partials producing stacking faults, making a computation of $\rho_d$ from lattice configurations much more difficult that the simple counting of near neighbour coordination which we did in 2D. The nucleation of stacking faults governed by their energy determines the dominant deformation mechanism in FCC crystals. Therefore considerable effort is needed to define and compute stacking fault energies\(^3\). Our results may offer an alternative. We have shown that the non-affine fraction $\phi$ provides an equally good measure for significant departure of the lattice from the topology of the ideal reference

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**Figure 10.** a. Plot of the non-affine fraction $\phi(T, \rho)$ for two temperatures $T = 0.35$ & 0.4 and at $\rho = 0.93$ & 0.99 for the 2DLJ solid. b. Data collapse plot of $\rho_d$, the defect density, plotted against the non-affine fraction $\phi$. Note that the data for $T = 0.4$ has also been plotted in Fig. 6b of the manuscript.

**Figure 11.** a–c Non affine particles in 3D-LJ, FCC solid of 16384 particles at $T=0.7$, $\rho=1.5$ for $\varepsilon = .02$, $\varepsilon = .04$ and $\varepsilon = .06$ respectively. Particles with less than 2 non-affine neighbours have been made translucent for clarity. Percolation occurs for $\varepsilon \gtrsim .04$ at this density where $\phi \approx 0.19$. 

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and may be used to understand pre-yielding phenomena in higher dimensional solids too. Of course, a computation of $\phi$ and $f_\phi$ needs data for instantaneous particle positions which may be available only in computer simulations or from colloidal solids.

Discussion

In summary, we have unearthed a hidden mechanical critical point associated with the percolation of non-affine droplets which is intimately tied to the onset of complex mechanical response in a crystalline solid. This finding is quite unexpected in a system as familiar as a crystalline solid. This transition does not manifest in mechanical or thermodynamic properties in the bulk, but subtly reveals itself in at least four ways:

(i) Dislocations, nucleated upon shear, are associated with a spatially extended non-affine cloud. The spatial scale is determined by two coarse-graining parameters $\chi_{\text{cut}}$ and $\Lambda$ (see Methods). It is this cloud of non-affinity that eventually percolates across the sample, the percolation transition being fairly robust to small changes in $\chi_{\text{cut}}$ and $\Lambda$.

(ii) The mechanical critical point is not apparent in bulk thermodynamics but in the restricted (finite size) thermodynamics of the localized droplets. The droplet fluctuations represent regions where the solid explores nearby minima in the free energy corresponding to metastable glassy or liquid configurations\(^2\)\(^8\). At an algorithmic level, since the droplets are small and transient, we require a special algorithm to distinguish them from the overwhelmingly large contribution of normal fluctuations in the equilibrium solid.

(iii) This mechanical critical point shows up in the distribution of local elastic stresses and local elastic constants and the emergence of strong spatial heterogeneity inaccessible to bulk elastic experiments.

(iv) Finally, the mechanical critical point is linked to the onset of a strong nonlinear dynamical response of the crystalline solid. Hysteretic response of the stress during strain cycling, slow creep relaxation of the strain under constant load and relaxation of stress below yield point are all manifestations of so-called “an-elastic” behaviour\(^2\)\(^9\) caused by non-linearities in the elastic response, elastic heterogeneities and time-dependent, slow, defect reorganisation\(^4\). While nonlinearities and anelasticity were long suspected to be connected with the appearance of lattice defects, we show that these features are precisely what characterises the hidden mechanical critical point associated with the percolation of non-affine droplets.

The emergence of plastic behaviour in amorphous solids appears to be associated with the percolation of localised non-affine deformations\(^3\)\(^0\). Indeed, a similar percolation transition for quasi-statically strained amorphous solids in the athermal ($T=0$) limit has been recently described in\(^3\)\(^1\). In the crystalline solid described here, percolation of non-affine droplets is, on the other hand, associated with the emergence of nonlinear, anelastic behaviour before the commencement of permanent plastic deformation.

Figure 12. a The percolation transition for non affine particles in the 3D-IJ, FCC solid at two values of $T=0.7$ & 0.8 and $\rho=1.5$ & 1.2. Inset shows the corresponding $\phi(\rho, T)$ curves. b Scaled stress-strain relations for the sheared solids are shown in a; symbols have the same meaning as in a. Non-affine particles percolate the solid at $\varepsilon/\varepsilon^*=1$. Inset shows the $\sigma$ vs $\varepsilon$ curve for $T=0.7$ and $\rho=1.5$. The arrow marks $\varepsilon^*$ and the solid line marks the linear Hooke’s law behaviour.
Nevertheless, it is tempting to hope that there may be an underlying common language describing the mechanical response of all solids (crystalline and amorphous) in general.

The hidden critical point betrays itself in our MD simulations only through local properties of non-affine droplets identified using carefully chosen cut-offs and thresholds. Before we end, we speculate on whether the critical point may be revealed by changing system parameters or by introducing novel forces so that it begins to affect bulk thermostrodynamics. We have shown in Ref. 12 that an external field \( h \), which couples to \( \chi \), may be tuned to control the equilibrium value of \( \chi \) and hence the defect density. This field may be introduced in MD simulations as well as realised experimentally in a colloidal solid using holographic optical tweezer techniques. The results presented here for \( h = 0 \) may be a reflection of an equilibrium phase transition in the full \( h - \sigma \) space occurring for \( h \neq 0 \). In the future, we would also like to study how such hidden critical points influence dynamical behaviour such as avalanches and intermittency.

Methods

Simulation details. Two dimensions. We prepare a triangular arrangement of 22,500 identical particles interacting pairwise via the LJ potential. Equilibration of our 2D model system is carried out at the fixed state point (fixed \( T \), \( \Omega \), and \( \rho \)) within the LAMMPS package (http://lammps.sandia.gov) and preserving both volume and number. Solids at each \( \rho \) and \( T \) are equilibrated with 10^8 molecular dynamics time-steps (MDS) where each time-step equals 10^{-4} LJ time units. Temperature fluctuations measured at equilibrium is of the order of 10^{-3}. Pure shear is then applied to the system following the protocol: \( L_x' = (1 + \varepsilon) L_x \) and \( L_y' = (1 - \varepsilon) L_y \), \( L_x \) and \( L_y \) denoting the box lengths in \( X \)- and \( Y \)-directions respectively with primed quantities representing the same after shear, \( \varepsilon \) is the shear step set to 0.002. After each shear step, the system is equilibrated for 4 \times 10^4 MDS. Configurations for the analysis is stored in the first 1 \times 10^4 MDS in regular intervals of 20 MDS. This cumulative shearing process is continued till the solid fails at the yield point \( \varepsilon_c \). The strain rate associated with the process therefore equals to 10^{-3} per LJ unit. The sound velocity propagating through the solid \( c_s = \sqrt{G/\rho} \) where \( G \) is the elastic modulus for pure shear and \( \rho \) is the density of the medium. For \( \rho = 0.99 \), \( L_x = 174.96 \) and \( L_y = 151.51 \) in LJ units), sound at velocity \( c_s \approx 6.12 \) in LJ units takes approximately 27 LJ time unit to travel across the simulation box in \( X \)-direction. Considering this, we refer to our shear protocol as quasi-static. The model solid at various densities is strained using the same quasi-static shear protocol.

Three dimensions. The number of particles is 16384 i.e. 16 × 16 × 16 cubic unit-cells of the FCC lattice, each containing 4 basis atoms. The same LJ interactions and the same ensemble with fixed number, volume and \( T \) (using the DPD thermostat) are used. The solid is in pure shear with \( L_x \) being extended by \( \varepsilon \), and \( L_y \) and \( L_z \) shortened such that the volume (and density) is preserved. The strain step is 0.002 as before. At every strain step the solid is equilibrated for 1.5 \times 10^5 MDS with a time step of 1. \times 10^{-3} in LJ units. Data is collected for the next 5. \times 10^4 at every 500 steps.

Identification of droplets. We identify non-affine droplets using the prescription described in. Briefly, at a fixed state point (fixed \( T \), \( \rho \) and \( \varepsilon \)), we calculate \( \chi \) for each particle within a neighbourhood \( \Omega \), defined by \( r < \Lambda \) for all configurations using the \( T = 0 \) defect free lattice at the same \( \rho \) and \( \varepsilon \) as reference. We next obtain the distribution \( P(\chi) \). We identify a fixed cutoff \( \chi_{cut} \) above which the fluctuations are deemed anharmonic. All particles in a single snapshot having \( \chi > \chi_{cut} \) are tagged and tagged particles residing within the first nearest neighbor shell of another tagged particle belong to the same cluster; any other cluster having at least 7 particles is identified as a non-affine droplet. Note that \( \Lambda \) and \( \chi_{cut} \) are parameters which are chosen such that \( \Lambda = 2.5 \) (2D) or 1.1 (3D) and \( \chi_{cut} \) excludes 90% of the weight of the first peak in \( P(\chi) \). Small deviations of these parameters do not change our results substantially.

Local thermodynamic quantities. We describe below our procedure for obtaining the local thermodynamic quantities for the droplets.

Local density \( \rho_c \). Once a droplet has been identified, we include one extra layer of particles so that the Voronoi nearest neighbors of all the particles within the cluster are included (whether non-affine or not). Then we count the number of Delaunay triangles using GEOMPACK (code available at http://people.sc.fsu.edu/~jburkardt/f_src/geompack1). The correct density of the droplet is then given by half of the ratio between number of particles \( n_i \) in the droplet and the total area of all the triangles \( \rho_c = n_i/(2A) \).

Local pressure \( p_c \). This is computed using the virial \( \langle F_{ii} \cdot r_{ij} \rangle \) where \( F_{ii} \) and \( r_{ij} \) are the nearest neighbor forces and distances respectively for particles \( i \) and \( j \) belonging to the droplet. The average \( \langle ... \rangle \) is over all \( n_i \). The droplets are characterized by a distribution of \( \rho_c \) and excess pressures, \( \Delta p_c = p - p_c \), where \( p \) is the mean pressure of the surrounding solid.
Block stress $\sigma_{B}$: To compute $\sigma_{B}$, we divide the simulation cell into sub-blocks $B$ of equal size. The stress is computed by averaging the virial for all particles $N_B$ in the blocks, over all the blocks and finally, over many independent configurations. The components of the stress distribution are calculated as 

$$
\sigma_{\alpha\gamma}^{(B)} = \frac{1}{2N_B} \sum_{k,l} (x_{k\alpha}^{(l)} - x_{k\alpha}^{(B)}) f_{\gamma}^{(l)}
$$

where $k,l$ are particle indices while $\alpha\gamma$ denotes the cartesian components of the stress.

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### Author Contributions

M.R. and S.S. conceived of the project and wrote the paper. T.D. did the necessary calculations, designed the specific algorithms and helped in writing the paper. S.G. checked the calculations and helped in designing algorithms and obtaining the data.

### Additional Information

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