Towards glasses with permanent stability

Taiki Yanagishima,1, 2 John Russo,3 Roel P. A. Dullens,1, 4 and Hajime Tanaka5, 6

1Department of Chemistry, Physical and Theoretical Chemistry Laboratory, University of Oxford, South Parks Road, OX1 3QZ, United Kingdom
2Department of Physics, Graduate School of Science, Kyoto University, Kitashirakawa Oiwake-cho, Sakyo-ku, Kyoto, 606-8502, Japan
3Department of Physics, Sapienza University of Rome, P. le Aldo Moro 5, 00185 Rome, Italy
4Institute for Molecules and Materials, Radboud University, Heyendaalseweg 135, 6525 AJ Nijmegen, The Netherlands
5Department of Fundamental Engineering, Institute of Industrial Science, University of Tokyo, 4-6-1 Komaba, Meguro-ku, Tokyo 153-8905, Japan
6Research Center for Advanced Science and Technology, University of Tokyo, 4-6-1 Komaba, Meguro-ku, Tokyo 153-8905, Japan

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Unlike crystals, glasses age or devitrify over time, reflecting their non-equilibrium nature. This lack of stability is a serious issue in many industrial applications. Here, we show by numerical simulations that the devitrification of quasi-hard-sphere glasses is prevented by suppressing volume-fraction inhomogeneities. A monodisperse glass known to devitrify with ‘avalanche’-like intermittent dynamics is subjected to small iterative adjustments to particle sizes to make the local volume fractions spatially uniform. We find that this entirely prevents structural relaxation and devitrification over aging time scales, even in the presence of crystallites. There is a dramatic homogenization in the number of load-bearing nearest neighbors each particle has, indicating that ultra-stable glasses may be formed via ‘mechanical homogenization’. Our finding provides a physical principle for glass stabilization and opens a novel route to the formation of mechanically stabilized glasses.

Glassy materials are known to spontaneously age and devitrify [1]. The gradual aging-driven drift in the physical properties of glasses over time is a serious issue for their applications. For example, localized crystallization can be detrimental to aqueous media in cryogenics [2–4], pharmaceuticals [5–7] and optical media [8–10]. There can be detrimental to aqueous media in cryogenics [2–4], and devitrify with ‘avalanche’-like intermittent dynamics [14–17]. Howroborated a relationship between local structure, local volume fraction [15], we take an orthogonal approach to previous works by removing inhomogeneities in local volume fractions and studying the subsequent dynamics. This strategy is closely linked to studies of hyperuniform packings, themselves characterized by suppressed long-range spatial correlations in local density [18, 19]. The system we study is a dense glass of particles interacting via a repulsive Weeks-Chandler-Andersen (WCA) potential at thermal energy $k_B T = 0.025$, propagated over time using standard Brownian Dynamics (overdamped Langevin dynamics) [20]. The system is commonly regarded as closely approximating hard spheres or experimental colloidal systems and has been widely used as an ideal model system to study aging and devitrification [14–17, 21–25]. Firstly, ‘conventional’ glassy (CG) states were prepared using a modified version of the Lubachevsky-Stillinger algorithm [26], identical to the method used in previous work [15]. Initial particle sizes are monodisperse, $\sigma_i = 1$ for all $i$, where $\sigma_i$ is the size of particle $i$ in the repulsive WCA potential. After an initial configuration is generated at some bulk volume fraction $\phi_0$, the energy of the packing is minimized using the FIRE algorithm [27] to relax residual stresses introduced by the quench.

In order to flatten the spatial volume fraction profiles, we adapt a recently developed method for making hyperuniform packings [19]. The local volume fraction, $\phi_i$, of particle $i$ is estimated using a radical Voronoi tessellation. Particles are then resized such that, for each Voronoi cell, $\pi \sigma_i^3/(6 v_i) = \phi_0$, where $v_i$ is the local Voronoi volume. The particle is then replaced in the center of the cell. As radical Voronoi cell boundaries change when the particle is resized, the state is no longer at an energy minimum. Thus, the packing is rerun through the FIRE algorithm, and the process repeated, checking that the standard deviation of local densities $\Delta \phi_i / \langle \phi_i \rangle$ decreases (see Fig. S1
This iterative process is stopped once changes to particle sizes no longer result in a narrowing of the local volume fraction distribution. We call this final glass state a “uniform glass (UG)” with its uniform local volume fraction over space. An illustration of the algorithm, the narrowing of the volume fraction profile, and the slight broadening of the particle size distribution are shown in Figs. 1(a), (b), and (c), respectively. In this paper, we express volume fractions using an effective volume fraction $\phi_{\text{eff}}$ with each iteration. (b) The probability density function of effective local volume fractions $\phi_{\text{eff}}^i$ with each iteration. (c) The probability density function of particle sizes $\sigma_i$ with each iteration.

To quantify the action of this $\phi_i$ flattening algorithm, we generate 50 independent CG/UG pairs. When $\phi_{\text{eff}}^0 = 0.68$, the initial CG glasses we generate have a $\phi_i$ distribution with a standard deviation $\Delta\phi_i/\langle\phi_i\rangle$ ranging from 3.7% to 3.9% over the 50 samples. After flattening, this is reduced to 0.37% to 0.95% (mean 0.63%), with a particle size polydispersity of 3.0% to 3.7% (mean 3.4%). Notably, this polydispersity is significantly lower than the dispersion required to prevent crystallization [28, 30–32]; i.e., one would still expect a strong thermodynamic driving force towards crystallization. We emphasize that this polydispersity approaches the one present in the most monodisperse colloidal suspensions used to experimentally study crystallization [23]. To show that the properties of the UG state are independent of polydispersity, we generate independent CG states that share the same particle size distribution of UG states, and refer to them as re-CG states in the following.

The fact that the UG states are prepared by suppressing local $\phi_i$ fluctuations suggests a natural connection between these states and hyperuniform configurations [18, 19, 34, 35], as mentioned above. To verify this, in Fig. 2 we plot the spectral density of the configurations $\chi_q$, defined as a power spectrum $\chi_q(q) = V^{-1}\langle I_q I_q^* \rangle$. $I_q$ is the Fourier transform of $I(r)$, the indicator function for a polydisperse suspension of hard spheres defined as $I(r) = \Sigma_i \Theta(|r - r_i| - R_i)$, where $r_i$ is the position of particle $i$, $R_i = \sigma_i/2$ is the radius, $\Theta$ is the Heaviside function, and $V$ is the total volume [34, 35]. In the context of an experiment, $\chi_q$ is simply the power spectrum of a binary ‘image’ of particles. It is clear that the $q \rightarrow 0$ limit of $\chi_q$ is diminished with progressive iterations. However, it is also clear that the $q^4$-scaling at low $q$ expected for class I hyperuniform systems [18] is not reached. We also note that the distribution of number densities $n_i$ significantly broadens; since the algorithm keeps the volume fraction constant, the size polydispersity results in a broader number density distribution, as shown in the inset of Fig. 2 Overall, it is clear that our UG states are approaching a hyperuniform state, but cannot be classed as hyperuniform themselves.

Now, we turn our attention to the dynamics. Brownian dynamics is applied to CG states, states reached after flattening [28], the initial CG glasses we generate have a $\phi_i$ distribution with a standard deviation $\Delta\phi_i/\langle\phi_i\rangle$ ranging from 3.7% to 3.9% over the 50 samples. After flattening, this is reduced to 0.37% to 0.95% (mean 0.63%), with a particle size polydispersity of 3.0% to 3.7% (mean 3.4%). Notably, this polydispersity is significantly lower than the dispersion required to prevent crystallization [28, 30–32]; i.e., one would still expect a strong thermodynamic driving force towards crystallization. We emphasize that this polydispersity approaches the one present in the most monodisperse colloidal suspensions used to experimentally study crystallization [23]. To show that the properties of the UG state are independent of polydispersity, we generate independent CG states that share the same particle size distribution of UG states, and refer to them as re-CG states in the following.

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\begin{figure}[h]
\centering
\includegraphics[width=\textwidth]{figure1.png}
\caption{Homogenization of local volume fractions. (a) Illustration of the $\phi_i$ flattening algorithm. A Voronoi tessellation is taken of an energy minimized configuration (solid blue). Particles are resized to match the local $\phi_i$ (dashed red) before being relocated to the center of the Voronoi cell (crosses). The energy is minimized again before the process is repeated until this global operation no longer reduces the standard deviation in the local volume fractions. (b) The probability density function of effective local volume fractions $\phi_{\text{eff}}^i$ with each iteration. (c) The probability density function of particle sizes $\sigma_i$ with each iteration.}
\end{figure}

\begin{figure}[h]
\centering
\includegraphics[width=\textwidth]{figure2.png}
\caption{Spectral density of static structures, from CG to UG. Spectral densities $\chi_q$ are given for progressive iterations as the CG state is transformed into a UG state. As the iterations progress, $\lim_{q \rightarrow 0} \chi_q(q)$ systematically decreases, though the scaling does not reach the $q^4$-scaling of a class I hyperuniform state. (inset) Distribution of local number densities $n_i$. The UG state features a significantly wider distribution of $n_i$.}
\end{figure}
FIG. 3. Impact of flattening $\phi_i$ over space on avalanche events and devitrification. (a) Squared displacements over ten $5000\tau_B$ runs initiated from the same CG state, the state after one iteration of the $\phi_i$ flattening algorithm, and the final UG state, as indicated. (b) Squared displacements over ten $5000\tau_B$ runs initiated from ten independently generated CG states using the same particle sizes as in the UG state (re-CG). Squared displacements for the UG state from (a) are also drawn for comparison. (c) Change in crystalline particles and force network for corresponding CG and UG states. The CG-generated state experiences growth in crystallinity, while the UG state created from it does not change. Changes in force connections between nearest neighbors are also shown. The thin blue lines indicate the original network; red and green lines at $t = 5000\tau_B$ indicate connections broken and formed.

The central question now becomes the nature of the avalanche-like dynamics. Figure S3 [28] shows crystalline particles in energy-minimized CG and UG states before and after dynamical propagation using Brownian dynamics. Crystallites are detected using nearest neighbor bond coherence in bond-orientational order parameter $q_6$, as in Refs. [36, 37]. Note that the UG states see no significant growth, as if the volume fraction flattening effectively ‘freezes’ post-critical nuclei. We stress that the bulk volume fraction $\phi_0$ and temperature remain unchanged. This is doubly clear by visualizing crystalline particles in CG and UG states over time as they thermally fluctuate (see Supplementary Movie [28]). Despite some fluctuations in bond coherence, the crystallinity of the UG state remains unchanged, while the CG state experiences significant crystal growth. This strongly corroborates previous findings that the avalanches provide the necessary perturbation for existing crystals to grow at deep supercooling [14, 15].

It is worth emphasizing that this ‘stability’ against avalanche-mediated crystallization is achieved by introducing only a minimal polydispersity in particle size. For the UG configurations used to calculate squared displacements in Figs. 3(a) and (b), this is 3.60%. Figure 3(b) compares the evolution of 10 UG states and their respective re-CG states (i.e., having the same size distribution but with an inhomogeneous local volume fraction distribution), showing that re-CG states are prone to avalanche displacements and that size polydispersity alone does not play a role in the stabilization of the UG states.

We now examine the structural changes that occur going from CG to UG states. At the level of pair correlations, we see that these states are virtually identical: Figure S2 [28] plots the radial distribution function $g(r)$ for CG, UG, and re-CG states, showing that the $g(r)$ of UG and re-CG states (which have the same particle-size distribution) are almost indistinguishable. Pair correlations thus are unable to capture the sharp discrepancy between the dynamics of UG and re-CG states. In Fig. 4(a), we plot the parameter $\langle Q_6 \rangle$, a coarse-grained measure of bond-orientational order $q_6$ averaged over nearest neighbors [35], and the energy per particle $\beta \langle U_i \rangle$ for 50 independently generated CG states, corresponding UG states and re-CG states generated from the particle size distribution of the UG states. While $\langle Q_6 \rangle$ fails to distinguish between (re-)CG and UG states, the UG states are found to be considerably lower in the energy landscape than their CG counterparts. Also the compressibility factor $Z = \beta P_V / (\sigma^3)$, where $P_V$ is the virial pressure, and the deviatoric stress invariant $J_2 = tr(s^2)/2$, where $s$ is the stress tensor, both plotted in Fig. 4(b), show a significant drop going from (re-)CG states to UG states. Note that the drop is greater over subsequent iterations of the $\phi_i$-flattening algorithm (see Fig. S3 [28]). We conclude that the system is annealed going from CG to UG states, with drops in both internal stress and energy, but this transformation is distinct from aging, as no significant changes in local structure occur at the pair (e.g., $g(r)$), and many-body level (e.g., $\langle Q_6 \rangle$). Thus, the UG state is not reached thermodynamically and should not be confused with the ideal glass, which is defined in a thermodynamic context [25, 29, 30, 40].

The central question now becomes the nature of the subtle change in structure that causes such a dramatic change in stability. To this end, we investigate the distribution of force neighbors, which were shown to describe
the onset of mechanical stability in glasses [15] and gels [12]. For particle $i$ with size $\sigma_i$, we measure the number of nearest neighbors $j$ which are located at a distance $r$ such that $r < 2^\alpha (\sigma_i + \sigma_j) \times \frac{1}{2}$, the interaction range of the WCA potential. These ‘force’ neighbors (FN) exert a repulsive force on particle $i$, and create linkages in the force chain network of the configuration; we let $n_{FN}$ be the number of force neighbors surrounding each particle. Figure 4(c) shows $(n_{FN})$ for particles with different local volume fractions $\phi_i^{\text{eff}}$ in a typical CG (red) state and its corresponding UG (blue) state. Distributions are given for both energy minimized (i.e., inherent) states and a thermally fluctuating configuration at the beginning of a Brownian Dynamics trajectory ($t = 5\tau_B$).

Firstly, we see that the CG state exhibits a linear relationship between the local volume fraction of a particle $\phi_i^{\text{eff}}$ and the average number of force neighbors it has, $(n_{FN}(\phi_i))$. With thermal fluctuations, the same trend is seen, except with fewer force connections. Now, going to the corresponding UG state, we find that the trend is closely preserved. The consequence of this is that the narrowing of the local volume fraction distribution directly results in a narrowing of the distribution of the number of force neighbor particles. The large error bars at the extrema are due to the small number of particles in the corresponding bins ($< 10$). The narrowing of the $n_{FN}$ distribution is further illustrated by considering changes in the number of force neighbors $\Delta n_{FN}$ for individual particles with different $\phi_i^{\text{eff}}$ in the CG state as it is transformed into a UG state (see inset of Fig. 4(c)). This confirms that particles at lower $\phi_i^{\text{eff}}$ gain force neighbors, while those with higher $\phi_i^{\text{eff}}$ lose them. Both these findings indicate that the narrowing of the distribution in $\phi_i^{\text{eff}}$ leads directly to a narrower, more homogeneous distribution of $n_{FN}$, i.e., mechanical homogenization. This strongly suggests that the mechanism behind the exceptional resistance to crystallization in the UG state is mechanical. Note that the same analysis of a re-CG state recovers the same broad range of $\phi_i^{\text{eff}}$ and the resulting range of $n_{FN}$ as the CG-state, corresponding to the recovery of avalanche dynamics. The same trends can be seen when CG and UG states are prepared at different volume fractions, as shown in Fig. S4 [25].

Local stresses are also affected by the homogenization of local connections. In Fig. 4 we plot both local hydrostatic (d) and deviatoric (e) pressures as a function of the number of force neighbors averaged over the same 50 independent configurations used for Figs. 4(a) and (b). In Fig. 4(d) we plot the average excess pressure contribution, $(P_{ex,i})$ for particles with different $n_{FN}$: in contrast to CG and re-CG states, whose excess pressures increase with $n_{FN}$, UG states have a maximum around $n_{FN} = 9$, close to the average $n_{FN}$ at the considered volume fraction. Similar behavior is found in the local deviatoric stress invariant $J_{2,i}$ plotted in Fig. 4(e), in that more local contacts $n_{FN}$ results in a lower deviatoric stress, though there is no maximum at $n_{FN} = 9$. The local arrangements of nearest neighbors in the UG state result in significant reductions in both volume (hydrostatic) and shape (deviatoric) altering stresses acting at the local level. In conclusion, the enhanced mechanical stability of the UG states is due to both a narrowing in the range of $n_{FN}$ and a reduction of local stresses.

The action of mechanical homogenization on devitrification may be understood as rendering configurations free of force chain network defects, making the system more resistant to thermal fluctuations. The connection between mechanical rigidity and glassy dynamics is consistent with recent work, which showed that a percolated force network is spontaneously formed below an experimental glass transition temperature, leading to the emergence of shear rigidity [11] [22]. Applied to our system, the destabilization of this percolated force network is the only mechanism by which our configurations may rearrange, i.e., triggering avalanches and causing structural...
aging and devitrification \[15\]; thus, by minimizing heterogeneities in the force network, we see a significant elevation in stability. The robustness that this imparts may be observed directly in Fig. 3(c) and the Supplementary Movie \[28\], which also shows force network connections broken (red), formed (green), and kept (blue) over time. It is worth noting that the combined simulation time of the Brownian Dynamics trajectories corresponds to at least 50,000$\tau_B$. For a typical colloidal experiment using 1-micron diameter particles at room temperature in water, this corresponds to 30 hours without a single rearrangement event, rendering our UG states ‘permanently’ stable within the observation time. We may also demonstrate this by considering perturbations to the UG state that artificially ‘trigger’ avalanche events in specific locations by removing or “voiding” single particles; due to the uniformly stable force network structure, avalanche events are highly localized, in sharp contrast to those seen for deeply supercooled conventional glasses \[14, 17\] (see \[28\] for details). All these results support the intrinsic link between the stability of uniform glasses and mechanical homogenization.

In summary, we successfully draw a correlation between the stability of a glass state and mechanical homogeneity. We believe this to be the first time avalanche dynamics has been directly related to a physical mechanism. Importantly, our results also pave the way towards developing a practical method to prepare mechanically stabilized homogeneous glasses. Experimental work \[43\] has shown that a colloidal system exhibits a transition to an absorbing state under oscillatory shear via so-called “random organization”, where the critical absorbing state is, in fact, hyperuniform \[44–47\]. Formation of a hyperuniform state in binary charged colloids has also been suggested \[48\]. Given that our UG states are ‘on the way’ to hyperuniformity, these recent studies suggest that any experimental efforts to approach such a state may readily realize mechanically stable UG states. Thus, we may realistically expect that our findings inspire experimental efforts not only to reach hyperuniformity, but to realize glasses that can resist aging. Such a method would significantly impact glass applications, as it would enable the preparation of ultra-stable glasses with unparalleled stability against degradation or devitrification over time.

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*tanaka@iis.u-tokyo.ac.jp

[1] C. Austin Angell, Kia L. Ngai, Greg B. McKenna, Paul F. McMillan, and Steve W. Martin, “Relaxation in glass-forming liquids and amorphous solids,” J. Appl. Phys. 88, 3113–3157 (2000).
[2] X. Han, H. B. Ma, C. Wilson, and J. K. Critser, “Effects of nanoparticles on the nucleation and devitrification temperatures of polyl cryoprotectant solutions,” Microfluid. Nanofluid. 4, 357–361 (2008).
[3] Charles J. Hunt, “Cryopreservation of human stem cells for clinical application: A review,” Transfus. Med. Hemother 38, 107–123 (2011).
[4] M. Sansinena, M. V. Santos, G. Taminelli, and N. Zaritzky, “Implications of storage and handling conditions on glass transition and potential devitrification of oocytes and embryos,” Theriogenology 82, 373–378 (2014).
[5] D. Q. M. Craig, P. G. Royall, V. L. Kett, and M. L. Hopton, “The relevance of the amorphous state to pharmaceutical dosage forms: glassy drugs and freeze dried systems,” Int. J. Pharm. 179, 179–207 (1999).
[6] J. Knapik-Kowalcuzk, Z. Wojnarowska, K. Chmiel, M. Rams-Baron, L. Tajber, and M. Paluch, “Can storage time improve the physical stability of amorphous pharmaceuticals with tautomeration ability exposed to compression? The case of a chloramphenicol drug,” Mol. Pharm. 15, 1928–1940 (2018).
[7] Nitin Salunkhe, Namdeo Jadhav, Harinhath More, and Prafulla Choudhari, “Sericin inhibits devitrification of amorphous drugs,” AAPS PharmSciTech 20, 285 (2019).
[8] M. J. Vestel, D. S. Grummon, R. Gronsky, and A. P. Pisano, “Effect of temperature on the devitrification kinetics of NiTi films,” Acta Mater. 51, 5309–5318 (2003).
[9] Y. Lin, M. M. Smedskjaer, and J. C. Mauro, “Structure, properties, and fabrication of calcium aluminate-based glasses,” Int. J. Appl. Glass Sci. 10, 488–501 (2019).
[10] M. Ordu, J. Guo, A. E. Akosman, S. Erramilli, S. Ramachandran, and S. N. Basu, “Effect of Thermal Annealing on Mid-Infrared Transmission in Semiconductor Alloy-Core Glass-Cladded Fibers,” Adv. Fiber Mat. 2, 178–184 (2020).
[11] A. L. Greer, “Partially or fully devitrified alloys for mechanical properties,” Mater. Sci. Eng., A 304–306, 68–72 (2001).
[12] J. Formell, S. González, E. Rossinyol, S. Suriñach, M. D. Baró, D. V. Louguine-Luzgin, J. H. Perezpekzo, J. Sort, and A. Inoue, “Enhanced mechanical properties due to structural changes induced by devitrification in Fe-Co-B-Si-Nb bulk metallic glass,” Acta Mater. 58, 6256–6266 (2010).
[13] D. V. Louzguine-Luzgin, “Vitrification and devitrification processes in metallic glasses,” J. Alloys Compd. 586, S2–S8 (2014).
[14] E. Sanz, C. Valeriani, E. Zaccarelli, W. C. K. Poon, Michael E. Cates, and P. N. Pusey, “Avalanches mediate crystallization in a hard-sphere glass.” Proc. Natl. Acad. Sci. 111, 75–80 (2014).
[15] T. Yanagishima, J. Russo, and H. Tanaka, “Common mechanism of thermodynamic and mechanical origin for aging and crystallization of glasses,” Nat. Commun. 8, 15954 (2017).
[16] N. B. Simeonova, R. P. A. Dullens, D. G. A. L. Aarts, V. W. A. de Villeneuve, H. N. W. Lekkerkerker, and W. K. Kegel, “Devitrification of colloidal glasses in real space,” Phys. Rev. E 73, 041401 (2006).
[17] D. Ganapathi, D. Chakrabarti, A. K. Sood, and R. Ganapathy, “Structure determines where crystallization occurs in a soft colloidal glass,” Nat. Phys. 17, 114–120 (2021).
[18] S. Torquato, “Hyperuniform states of matter,” Phys. Rep. 745, 1–95 (2018).
[19] J. Kim and S. Torquato, “Methodology to construct large realizations of perfectly hyperuniform disordered packings,” Phys. Rev. E 99, 052141 (2019).
[20] D. L. Ernack and J. A. McCammon, “Brownian dynamics with hydrodynamic interactions,” J. Chem. Phys. 69, 1352–1360 (1978).
[21] V. A. Martinez, G. Bryant, and W. van Megen, “Slow dynamics and aging of a colloidal hard sphere glass,” Phys. Rev. Lett. 101, 135702 (2008).
[22] Jennifer M. Lynch, Gianguido C. Cianci, and Eric R. Weeks, “Dynamics and structure of an aging binary colloidal glass,” Phys. Rev. E 78, 031410 (2008).
[23] E. Zaccarelli, C. Valeriani, E. Sanz, W. C. K. Poon, M. E. Cates, and P. N. Pusey, “Crystallization of hard-sphere glasses,” Phys. Rev. Lett. 103, 135704 (2009).
[24] Rojman Zargar, Bernard Nienhuis, Peter Schall, and Daniel Bonn, “Direct measurement of the free energy of aging hard sphere colloidal glasses,” Phys. Rev. Lett. 110, 255301 (2013).
[25] Takeshi Kawasaki and Hajime Tanaka, “Structural evolution in the aging process of supercooled colloidal liquids,” Phys. Rev. E 89, 062315 (2014).
[26] B. D. Lubachevsky and F. H. Stillinger, “Geometric properties of random disk packings,” J. Stat. Phys 60, 561–583 (1990).
[27] E. Bitzek, P. Koskinen, F. Gähler, M. Moseler, and P. Gumbsch, “Structural relaxation made simple,” Phys. Rev. Lett. 97, 170201 (2006).
[28] See Supplemental Material for information on the UG algorithm, radial distribution function, compressibility over iterations, UG glasses at different volume fractions, artificially triggered avalanches, and Supplementary Movie.
[29] D. M. Heyes and H. Okumura, “Equation of state and structural properties of the Weeks-Chandler-Andersen fluid,” J. Chem. Phys. 124 (2006), 10.1063/1.2176675.
[30] I. Moriguchi, K. Kawasaki, and T. Kawalaatsu, “The effects of size polydispersity in nearly hard sphere colloids,” J. Phys. II 3, 1179–1184 (1993).
[31] P. Bartlett, “A geometrically-based mean-field theory of polydisperse hard-sphere mixtures,” J. Chem. Phys. 107, 188–196 (1997).
[32] Hajime Tanaka, Takeshi Kawasaki, Hiroshi Shintani, and Keiji Watanabe, “Critical-like behaviour of glass-forming liquids,” Nat. Mater. 9, 324–331 (2010).
[33] T. Palberg, “Crystallization kinetics of colloidal model suspensions: recent achievements and new perspectives.” J. Phys.: Condens. Matter 26, 333101 (2014).
[34] C. E. Zachary, Y. Jiao, and S. Torquato, “Hyperuniform long-range correlations are a signature of disordered jammed hard-particle packings,” Phys. Rev. Lett. 106, 178001 (2011).
[35] C. E. Zachary, Y. Jiao, and S. Torquato, “Hyperuniformity, quasi-long-range correlations, and void-space constraints in maximally random jammed particle packings. I. Polydisperse spheres,” Phys. Rev. E 83, 051308 (2011).
[36] P. J. Steinhardt, D. R. Nelson, and M. Ronchetti, “Bond-orientational order in liquids and glasses,” Phys. Rev. B 28, 784–805 (1983).
[37] J. Russo and H. Tanaka, “The microscopic pathway to crystallization in supercooled liquids.” Sci. Rep. 2, 505 (2012).
[38] Wolfgang Lechner and Christoph Dellago, “Accurate determination of crystal structures based on averaged local bond order parameters,” J. Chem. Phys. 129, 114707 (2008).
[39] Andrea Cavagna, “Supercooled liquids for pedestrians,” Phys. Rep. 476, 51–124 (2009), 0903.4264.
[40] Hajime Tanaka, Hua Tong, Rui Shi, and John Russo, “Revealing key structural features hidden in liquids and glasses,” Nat. Rev. Phys. 1, 333–348 (2019).
[41] Hua Tong, Shiladitya Sengupta, and Hajime Tanaka, “Emergent solidity of amorphous materials as a consequence of mechanical self-organisation,” Nat. Commun. 11, 4863 (2020).
[42] Hideyo Tsurusawa, Mathieu Leocmach, John Russo, and Hajime Tanaka, “Direct link between mechanical stability in gels and percolation of isostatic particles,” Sci. Adv. 5, eaav6090 (2019).
[43] Laurent Corte, Paul M. Chaikin, Jerry P. Gollub, and David J. Pine, “Random organization in periodically driven systems,” Nat. Phys. 4, 420–424 (2008).
[44] Daniel Hexner and Dov Levine, “Hyperuniformity of critical absorbing states,” Phys. Rev. Lett. 114, 110602 (2015).
[45] Elsen Tjhung and Ludovic Berthier, “Hyperuniform density fluctuations and diverging dynamic correlations in periodically driven colloidal suspensions,” Phys. Rev. Lett. 114, 148301 (2015).
[46] Joost H. Weijls, Raphaël Jeanneret, Rémi Dreyfus, and Denis Bartolo, “Emergent hyperuniformity in periodically driven emulsions,” Phys. Rev. Lett. 115, 108301 (2015).
[47] Sam Wilken, Rodrigo E. Guerra, David J. Pine, and Paul M. Chaikin, “Hyperuniform structures formed by shearing colloidal suspensions,” Phys. Rev. Lett. 125, 148001 (2020).
[48] Duyu Chen, Enrique Lomba, and Salvatore Torquato, “Binary mixtures of charged colloids: a potential route to synthesize disordered hyperuniform materials,” Phys. Chem. Chem. Phys. 20, 17557–17562 (2018).