Hopping and the Stokes–Einstein relation breakdown in simple glass formers

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One of the most actively debated issues in the study of the glass transition is whether a mean-field description is a reasonable starting point for understanding experimental glass formers. Although the mean-field theory of the glass transition—like that of other statistical systems—is exact when the spatial dimension $d \to \infty$, the evolution of systems properties with $d$ may not be smooth. Finite-dimensional effects could dramatically change what happens in physical dimensions, $d = 2,3$. For standard phase transitions finite-dimensional effects are typically captured by renormalization group methods, but for glasses the corrections are much more subtle and only partially understood. Here, we investigate hopping between localized cages formed by neighboring particles in a model that allows to cleanly isolate that effect. By bringing together results from replica theory, cavity reconstruction, void percolation, and molecular dynamics, we obtain insights into how hopping induces a breakdown of the Stokes–Einstein relation and modifies the mean-field scenario in experimental systems. Although hopping is found to supersede the dynamical glass transition, it nonetheless leaves a sizable part of the critical regime untouched. By providing a constructive framework for identifying and quantifying the role of hopping, we thus take an important step toward describing dynamic facilitation in the framework of the mean-field theory of glasses.

Glasses are amorphous materials whose rigidity emerges from the mutual caging of their constituent particles—be they atoms, molecules, colloids, grains, or cells. Although glasses are ubiquitous, the microscopic description of their formation, rheology, and other dynamical features is still far from satisfying. Developing a more complete theoretical framework would not only resolve epistemological wrinkles (1), but also improve our material control and design capabilities. However, such a research program remains fraught with challenges. Conventional paradigms based on perturbative expansions around the low-density, ideal gas limit (for moderately dense gases and liquids) or on harmonic expansions around an ideal lattice (for crystals) fail badly. Because dense amorphous materials interact strongly, low-density expansions are unreliable, whereas harmonic expansions lack reference equilibrium particle positions. These fundamental difficulties must somehow be surmounted to describe the dynamical processes at play in glass formation.

A celebrated strategy for studying phase transitions is to consider first their mean-field description, which becomes exact when the spatial dimension $d$ of the system goes to infinity (2), before including corrections to this description. In that spirit, we open the $d \to \infty$ “ideal” random first-order transition (rRFOT) scenario, which, based on the analysis of simple models, brings together static- (3–5) and dynamics-based (mode-coupling) (6) results for glass formation (reviews in refs. 7 and 8) (8–11). In rRFOT, an infinitely slowly cooled simple liquid (or compressed hard sphere fluid) becomes infinitely viscous, i.e., forms a glass in which particles are completely caged, at the (critical) dynamical transition temperature $T_d$ (or packing fraction $\phi_d$). Upon approaching this transition, caging makes the diffusivity $D$ vanish as a power-law $D \sim (T - T_d)^\nu$, and the viscosity diverges as $\eta \sim (T - T_d)^{\eta_1}$. Hence, in the critical regime one expects the Stokes–Einstein relation (SER) between transport coefficients, $D \sim \eta^{1/\nu}$, to hold. In short, the $d \to \infty$ scenario is characterized by (i) a sharp dynamical glass transition associated with perfect caging, (ii) a power-law divergence of $\eta$, and (iii) the SER being obeyed.

As observed in ref. 12, the phenomenology of finite-dimensional systems is, however, quite different from the rRFOT scenario. In particular, it does not recapitulate elementary experimental observations, such as Vogel–Tammann–Fulcher (VTF) viscosity scaling in fragile glasses, $\eta \sim e^{\text{const.}/(T - T_0)}$ ($B_{\text{VTF}}$ and $T_0$ are phenomenological constants), and breakdown of the SER, $D \sim \eta^{1/\nu}$ (phenomenologically $\nu \gg 1$) (13–16). As a result, the relevance of the rRFOT picture for experimental systems remains the object of lively debates.

Part of the difficulty of clarifying the situation in finite $d$, where the rRFOT description is only approximate and the dynamical transition is but a crossover, lies in the sheer number of different contributions one has to take into account. From a purely field-theoretic point of view, one has to include finite-dimensional corrections to critical fluctuations. A Ginzburg criterion gives $d_a = 8$ as the upper critical dimension for the dynamical transition (17–20), and hence for $d < d_a$ critical fluctuations renormalize the power-law scaling exponents. In principle, these corrections could be captured by a perturbative $d_a - d$ expansion, and phenomenological arguments along this direction indicate that they could also induce a SER breakdown (17). A number of non-perturbative processes in $1/d$ must additionally be considered: (i) In the rRFOT picture, caging is perfect, and hence in the glass phase each particle is forever confined to a finite region of space delimited by its neighbors (6). However, it has been theoretically proved (21) and experimentally observed (22) that in low-dimensional systems the diffusivity is never strictly zero. Single particles can indeed hop between neighboring cages (23–26),

Significance

Like crystals, glasses are rigid because of the self-caging of their constituent particles. The key difference is that crystal formation is a sharp first-order phase transition at which cages form abruptly and remain stable, whereas glass formation entails the progressive emergence of cages. This loose caging complicates the description of the glass transition. In particular, an important transport mechanism in this regime, hopping, has thus far been difficult to characterize. Here we develop a completely microscopic description of hopping, which allows us to clearly assess its impact on transport anomalies, such as the breakdown of the Stokes–Einstein relation.

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and the free space they leave behind can facilitate the hopping of neighboring particles. Facilitation can thus result in cooperative hopping and avalanche formation (27–29). (ii) For some glass formers, activated crystal nucleation cannot be neglected and interferes with the dynamical arrest, leading to a glass composed of microscopic geometrically frustrated crystal domains (30). (iii) In the iRFOT scenario, the dynamical arrest is related to the emergence of a huge number of distinct metastable glass states whose lifetime is infinite. In finite dimensions, however, a complex glass–glass nucleation process gives a finite lifetime to these metastable states (5, 12, 31). The dynamics of glass-forming liquids are then profoundly affected. Including glass–glass nucleation into iRFOT leads to the complete RFO process (12), in which the mean-field dynamical glass transition becomes a crossover (12), and both the VTF scaling and facilitation can be recovered (32, 33).

Because the treatment of these different processes has thus far been mostly qualitative, their relative importance cannot be easily evaluated. A controlled first-principle, quantitative treatment is possible for the moment limited to the exact solution for $d \to \infty$ (10, 11, 34, 35). Its approximate extension to finite $d$ (6, 8, 36) completely ignores the nonperturbative effects mentioned above. This approach therefore cannot, on its own, cleanly disentangle the various corrections. Systematic studies of glass formation as a function of $d$ have encouragingly shown that these corrections are limited even down to $d = 3$ (15, 16, 37–40), provided length and timescales are not too large, as is typical of numerical simulations and experiments with colloids and grains. In particular, with increasing $d$ the distribution of particle displacements (the self-van Hove function) loses its second peak associated with hopping (16), the critical power-law regimes lengthen (41), and the SER breakdown weakens (15, 16, 40), which motivates investigating corrections to iRFOT in a controlled way.

Here we develop a way to isolate the simplest of these corrections, i.e., hopping, by studying a finite-dimensional mean-field model. Through the use of the cavity reconstruction methodology developed in the context of spin glass and information theory (42), we carefully describe caging, using self-consistent equations that can be solved numerically. We can thus compute the cage width distribution and isolate hopping processes. Our results provide an unprecedentedly clear view of the impact of hopping on the dynamical transition and on the SER breakdown in simple glass formers.

**MK Model**

We consider the infinite-range variant of the hard sphere(s) (HS)-based model proposed by Mari and Kurchan (MK) for simple structural glass formers (43–45) (details in *SI Text, section I.A*). The key feature of the MK model is that, even though each sphere has the same diameter $\sigma$, pairs of spheres interact via an additional constant shift that is randomly selected over the full system volume. This explicit quenched disorder eliminates the possibility of a crystal state, suppresses coherent activated barrier crossing that leads to glass–glass nucleation (44), and diminishes the possibility of facilitated hopping (as we discuss below). However, at finite densities the number of neighbors that interact with a given particle is finite and therefore finite-dimensional corrections related to hopping remain, in principle, possible.

MK liquids have a trivial structure. Even in the dense, strongly interacting regime, the pair correlation in the liquid phase is simply $g_2(r) = \Theta(r - \sigma)$ [where $\Theta(x)$ is the Heaviside step function], because particles are randomly displaced in space. In addition, even if both particles $i$ and $k$ are nearby particle $j$, they need not be close neighbors, and hence all higher-order structural correlations are perfectly factorizable. Because only two-body correlations contribute, the virial series can be truncated at the second virial coefficient (44), and hence the equation of state for pressure is trivially $pV^2/\rho = V_B$, where $V_B = V_2(1)/\sigma^d/2$ is the second-virial coefficient for $d$-dimensional hard spheres, $V_2(R)$ is the volume of a $d$-dimensional ball of radius $R$, $\rho$ is the number density [the packing fraction $\rho = \rho_V(\sigma/2)$], and the inverse temperature $\beta$ is set to unity (43–45) (*SI Text, section I.A*). Note that these structural features hold for the liquid phase of the MK model in all $d$ and for standard HS liquids in the limit $d \to \infty$ (8, 46). The MK model therefore coincides with standard HS in that limit. For a given finite $d$, however, MK liquids are structurally more similar to their $d \to \infty$ counterparts than HS liquids are. One thus sidesteps having to take into account the nontrivial structure of $g_2(r)$, which muddles the description of standard finite-dimensional HS (8).

For the MK model, one can easily construct equilibrated liquid configurations at all $\phi$, even for $\phi > \phi_g$. [For standard HS, by contrast, prohibitively long molecular dynamics (MD) simulations are necessary in this regime.] This dramatic speedup is accomplished by adapting the planting technique developed in the context of information theory (47) (*SI Text, section I.B*). It is thus possible to study MK liquids arbitrarily close to, both above and below, the dynamical glass transition at $\phi_g$. A systematic study of caging beyond $\phi_g$ is also possible due to the cavity reconstruction formalism (*SI Text, section I.A*), a method adapted from the statistical physics of random networks (42).

**Caging**

The MK model dynamics are studied by event-driven MD simulations of planted initial configurations with $N = 4,000$ particles (details in *SI Text, section I.B*) (37, 38). The mean square displacement (MSD) $\Delta(t) = \langle \sum_{i=1}^N |r_i(t) - r_i(0)|^2 \rangle/N$ is determined from time evolution of the particle positions $r_i(t)$. At short times, before any collision occurs, ballistic motion gives $\Delta(t) \propto d^2t^2$; at long times, diffusive motion gives $\Delta(t) \approx 2Dd^2 t$. From $\tau_{\text{ser}}$ onward, the ballistic and the diffusive regimes are separated by an intermediate cage regime in which $\Delta(t) \approx \Delta$ is approximately constant, first appearing as an inflection point and then as a flat-fledged plateau (Definition in *SI Text, section I.C*). Simply put, after a few collisions with its neighbors, a particle becomes confined to a small region of space of linear size $\sqrt{\Delta}$, from which it can escape, and henceforward diffuse, only after a very large number of collisions.

In the $d \to \infty$ iRFOT scenario, a sharp dynamical transition occurs at $\phi_g$ (6, 8, 10), beyond which complete caging results in an infinitely long plateau and in the disappearance of the diffusive regime. In finite-dimensional systems, one can use an approximate theory based on a Gaussian assumption for the cage shape, to obtain a prediction for $\phi_g$ and $\Delta$ (8, 45) (*SI Text, section II.A*). One can also estimate $\phi_g$ from the simulation results by fitting the diffusivity, using the mean-field critical form $D = \phi_g - \phi_g^2$, and $\Delta = \lim_{\tau \to \infty} \Delta(t)$ beyond $\phi_g$ (Fig. 1A). As expected from the suppression of various finite $d$ corrections, the critical power-law regime is much longer for the MK model than for standard finite-dimensional HS (Fig. 1B) (44). Marked qualitative discrepancies from the iRFOT predictions are nonetheless observed: (i) Numerical estimates for $\phi_g$ systematically deviate from the approximate Gaussian result for $\phi_g$ (Fig. 1C), even though the two quantities grow closer with dimension. (ii) The diffusion time $\tau_D \propto \sigma^2$ [Definition in *SI Text, section I.C*] and the structural relaxation time $\tau_{\text{ser}}$ (definitions and a discussion of this point in *SI Text, section I.C*) follow the SER, $\tau_D \propto \tau_{\text{ser}}$, from $\phi_{\text{ser}} \approx \phi_g$ to $\phi_{\text{ser}} < \phi_g$, but then the SER breaks down, $\tau_D \propto \tau_{\text{ser}}^2$ with $\omega = 0.22$, in all $d$ (Fig. 1D). With increasing $d$, however, the timescale for this crossover, $\tau_D(\phi_{\text{ser}})$, also increases (Fig. 1D), and thus $\phi_{\text{ser}}$ grows closer to $\phi_g$ and $\phi_g$ (Fig. 1C). (iii) Even above $\phi_g$, a steady drift of the MSD plateau can be detected (Fig. 1B), but the magnitude of this effect diminishes with increasing $d$.

To clarify the physical origin of the above discrepancies, we first determine whether the mismatch between $\phi_g$ and $\phi_{\text{ser}}$ is due to the hypothesis made in computing the latter, i.e., that all of the cages have a Gaussian shape of a fixed diameter $\Delta$, by using
Fig. 1. (A) MSD of the MK model in d = 3 for φ = 0.40, 1.00, 1.40, 1.65, 1.72, 1.78, 1.84, 1.93, 2.00, 2.20, and 2.50, from top to bottom. The onset of caging φ\textsubscript{onset} (red), the theoretical dynamical transition φ\textsubscript{dyt} (blue), and its dynamical estimate φ\textsubscript{dyt} (magenta) are highlighted. Note that at φ\textsubscript{dyt} and beyond a steady drift of the MSD plateau can be detected. (B) Power-law scaling in d = 3 of the characteristic time η\textsubscript{0} determined by fitting φ\textsubscript{dyt} = 1.93 and φ = 4.95 and by using the idealized mean-field result φ\textsubscript{dyt} = 1.78 and by fitting φ = 3.27. (Inset) Dimensional evolution of η\textsubscript{0} and η. The dashed line indicates the d = w result, η\textsubscript{0} = 2.3786 (34). Solid lines are guides for the eye. (C) The dimensional scaling of φ\textsubscript{dyt} and φ\textsubscript{dyt} converge as d increases, whereas the onset of caging at φ\textsubscript{onset} remains clearly distinct. The dashed line is the replica result φ\textsubscript{dyt} = 4.802\pm0.8 (8, 10). Solid lines are guides for the eye. (D) Dimensional re-scaling of the SER (black) and SER breakdown (red) regimes for the MK model with w = 0.22. (Inset) The ratio t\textsubscript{onset}/t\textsubscript{0} grows exponentially with d (solid line), where t\textsubscript{onset} = η\textsubscript{0}/φ\textsubscript{dyt} and t\textsubscript{0} is the microscopic time, i.e., the characteristic time for the decay of the velocity autocorrelation function (48) (details in SI Text, section I.C.2).

the cavity reconstruction formalism to relax both assumptions (42). Above φ\textsubscript{onset}, we can build the equilibrated neighborhood of particle i to self-consistently determine the overall cage size and/or shape distribution P(Δ) (details in SI Text, section II.A). The process involves placing Poisson-distributed neighbors j that are randomly assigned a cage size Δ\textsubscript{j} from a prior guess of P(Δ), with a fixed function shape f\textsubscript{A} (r) (a Gaussian or a ball function, for instance). Averaging over the vibrational relaxation of each neighboring particle gives the cavity field ψ(r) felt by particle i, which is the probability density of the particle being at position r (Fig. 2A). The existence of a cage centered around i is guaranteed by the cavity reconstruction procedure. The variance $\langle \Delta^2 \rangle = \langle r^2 \rangle - \langle r \rangle^2$ associated with the evolution of particle i within this cage, which can be computed through simple Monte Carlo sampling, provides the posterior caging radius Δ. Sufficient repeats of this determination provide a new estimate of $\bar{P}(\Delta)$, and iterating the overall procedure eventually converges to a fixed-point distribution $P_\infty(\Delta)$. We find that both Gaussian and ball caging functions give the same size distribution $P_\infty(\Delta)$ (Fig. 2B) and that $P_\infty(\Delta)$ is reasonably well approximated by a gamma distribution for all $\phi > \phi_\text{dyt}$ (Fig. 2C). The average cage size $\bar{\Delta}$ also quantitatively agrees with the analytical predictions of refs. 8 and 45 (Fig. 2D), including its characteristic square-root singularity upon approaching $\phi_\text{dyt}$, i.e., $\bar{\Delta}(\phi) \sim \sqrt{\phi - \phi_\text{dyt}}$. We thus conclude that the theoretical prediction of $\bar{\Delta}$ and $\phi_\text{dyt}$ is fairly insensitive to both the caging form and the second (or higher) moments of the cage size distribution, as well as to the theoretical method chosen (SI Text, section II.A).

It follows that deviations from the $d \rightarrow \infty$ scenario ought to be ascribed to an imperfect caging above $\phi_\text{dyt}$ in finite-dimensional systems. Microscopically, these imperfections correspond to particles trapped for a finite time before escaping to another cage through a narrow passage (Fig. 3A). Because the above calculations solely consider single-cage forms, a fixed-point distribution $P_\infty(\Delta)$ can only be reached by removing these “hopping” segments of the particle trajectories (details in SI Text, section II.A). Not only does $\phi_\text{dyt}$ then appear at higher densities, but as long as the network of connected cages percolates, dynamical arrest is also formally possible. In this context, it is interesting to note that for a prior $P_\infty(\Delta) = \delta(\Delta)$, the first iteration of the cavity reconstruction formalism is analogous to the void (Swiss-cheese) percolation setup for a Poisson process (49). In addition, for a nontrivial distribution of cage sizes, thresholding volume exclusion maps cavity reconstruction onto void percolation for polydisperse spheres (50) (SI Text, section II.C). This equivalence between cavity reconstruction and void percolation sheds light on the single-cage assumption. In the iRFOT description, the MSD of each particle should remain finite when $\phi > \phi_\text{dyt}$, but by construction the MSD can be truly bounded only if (minimally) $\phi > \phi_\text{dyt}$, the void percolation transition.

From MD simulations of the MK model, we detect the first hopping event of each particle (details in SI Text, section III.A). Around $\phi_\text{dyt}$, mode-coupling and hopping processes mix, but hopping quickly dominates the dynamics upon increasing $\phi$. Although the hopping of a particle does not leave an empty void in the MK model, it can nonetheless unblock a channel for a neighboring particle to leave its cage and hence facilitate its hopping. Facilitation is thus present, but weaker than in standard finite-dimensional HS, especially at high densities. Weakened
facilitation is notably signaled by the fact that the distribution of hopping times computed from a regular MD simulation largely coincides with the distribution obtained in the cavity procedure, where a single particle hops in an environment where neighboring particles are forbidden to do so (Fig. 3B, Inset). We find the cumulative distribution of hopping times over the accessible dynamical range to be well described by a power law $G_h(t) = (t/t_h)^{-\gamma}$ (Fig. 3B), with the characteristic hopping time $t_h$ increasing roughly exponentially with $\phi$ and markedly increasing with $d$ (Fig. 3D). This Arrhenius-like scaling form is consistent with a gradual and uncorrelated narrowing of the hopping channels with $\phi$. Note that similar phenomenological power-law distributions have recently been reported for other glass-forming systems, such as the bead-spring model for polymer chains (51). We get back to this point in section III.A.

**Finite-Dimensional Phase Diagram**

A clear scenario for hopping in the MK model follows from this analysis (Fig. 4). Dynamically, the system becomes increasingly sluggish upon increasing $\phi$ above $\phi_{\text{onset}}$. Initially, cages are not well formed and the slowdown exhibits a power-law scaling, according to the iRFOT critical predictions. Hopping cannot be defined because cages are too loose. Upon approaching $\phi_{\text{onset}}$, however, cages become much longer lived. In this regime, iRFOT predictions give a rapidly growing $t_h$, but beyond $\phi_{\text{onset}}$ hopping causes large deviations from this scaling. The critical-like behavior of the diffusivity is also pushed to denser systems, and fitting to a power law gives $\tau_h > \phi_{\text{onset}}$. When $t_h$ is comparable to $t_h$, a mixed regime emerges, characterized by a SER breakdown, as we discuss below. Even beyond $\phi_{\text{onset}}$, however, the dynamics are not fully arrested. Hopping remains possible, which shows that $\phi_{\text{onset}}$ has no fundamental meaning and is just a fitting parameter associated to an effective power-law divergence of $t_h$. In fact, the MK dynamical data are better fitted by a VTF form than by the critical power law (Fig. 4A), although the fitting parameter $\phi_{\text{onset}}$ has no direct static interpretation because it is intermediate between $\phi_d$ and $\phi_p$.

The dynamics can also be understood from the organization of cages. The critical density $\phi_d$ of iRFOT corresponds to the emergence of a connected network of cages. Typical networks for $\phi_d < \phi < \phi_p$ span the system volume. When $\phi > \phi_p$, they become finite and the mean network volume $V_{\text{net}}$ (sum of cage volumes in the network) follows a critical scaling from standard percolation (Fig. 4B). Based on this analysis, in the absence of facilitation the dynamical arrest should take place at $\phi_d$ (53). Note that although above $\phi_d$, the single-particle MSD is bounded, a particle can still explore a finite number of cages. Perfect single-cage trapping can be found at $\phi \to \infty$ only in finite $d$. Hopping is then infinitely suppressed because both the width and the number of hopping channels between cages vanish. However, even if hopping interferes with caging, well above $\phi_d$ vibrational relaxation within the cage is sufficiently quick to numerically distinguish it from hopping. This large separation of time-scales enables the facile detection of hopping in MD simulations and cavity reconstruction. However, upon approaching $\phi_d$ the task becomes acutely sensitive to the arbitrary thresholding inherent to any hopping detection algorithm (22, 54) (details in SI Text, section III.A).

As expected from the exactness of the iRFOT description in $d \to \infty$, $\phi_d/\phi_p \to 1$ with increasing $d$. Both $\gamma$ and $\beta$ also appear to converge to the $d = \infty$ value (Fig. 1B) (34). Because $\phi_d < \phi_p$ for all $d$, the suppression of hopping with increasing $d$ (Fig. 1D) Inset ought to be ascribed either to the narrowing of the hopping channels or to topological changes of the cage network. Because the pressure at the dynamical transition increases only slowly with

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**Fig. 3.** (A) Illustration of a cavity reconstruction in $d = 2$ for a hopping particle. In this case the neighboring particles allow the central particle to hop to other cages (red line). (B) Cumulative time probability distribution of hopping events $G_h(t)$ for $d = 3$ systems at densities (from top to bottom) $\phi = 1.76$, 1.84, 1.90, 1.97, and 2.10, along with the power-law scaling form (dashed line). (Inset) Single-particle hopping from the cavity reconstruction (circles) overlays with the MD simulations at short times ($\phi = 1.90$). (C and D) Phenomenological scaling parameters $\gamma$ and $\beta$ (as a guide for the eye for $\mu$ and exponential fits for $G_h$), which show that $\phi_d$ has no fundamental meaning and is just a fitting parameter associated to an effective power-law divergence of $t_h$. In fact, the MK dynamical data are better fitted by a VTF form than by the critical power law (Fig. 4A), although the fitting parameter $\phi_{\text{onset}}$ has no direct static interpretation because it is intermediate between $\phi_d$ and $\phi_p$.

**Fig. 4.** (A and B) Dynamical (A) and static phase (B) diagrams for the MK model in $d = 3$. Early in the critical regime, the relaxation times scale like a power law, but beyond $\phi_{\text{onset}}$, hopping causes large deviations from this scaling. An effective $\phi_h$ is numerically detected instead. A VTF scaling fits the data even better. Statically, cages can be detected from $\phi_h$ onward by removing hopping. In reality, the fine intercage channels that allow hopping result in a cage network. Beyond $\phi_d$, the typical network stops percolating and the network volume scales critically, $V_{\text{net}} \sim (\phi_p - \phi)^{-\beta}$ with $\beta = 2.40$ (dashed blue line) (49, 52). The single-cage limit is reached when $V_{\text{net}} \sim \phi_d^{-\beta}$. 

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dimension ($p_d \sim d$), the typical channel width is expected to stay roughly constant. The topology of the cage network, however, has a larger dimensional dependence. The cage network at percolation, for instance, has a fractal dimension $d_\text{f} \ll d$ (52); e.g., $d_\text{f} \approx 4$ for $d \geq d_\text{a} = 6$. Although this result is valid only at $\varphi_c$ proper, the local network structure persists at smaller $\varphi_c$ because the loss of the cage network fractality takes place through the single-point inclusion of nonpercolating clusters (52). The network topology is therefore such that the hopping channels (even assuming that their cross section remains constant) cover a vanishingly small fraction of the cage surface as $d$ increases. The limited number of ways out of a local cage thus entropically suppresses hopping.

### SER Breakdown

With hopping events clearly identified, it becomes possible to isolate the pure critical iRFOT (or mode-coupling) regime. Within this regime, we obtain a power-law scaling that is consistent with $\varphi_c$, $\varphi_u$, and $\varphi_{\text{SER}}$ converges as $d$ increases, whereas $\varphi_{\text{cset}}$ remains distinctly smaller (compare with Fig. 1C). Note that in $d=8$, $\varphi_u$, $\varphi_d$, and $\varphi_{\text{SER}}$ are numerically indistinguishable. (Inset) Dimensional evolution of $\gamma$ and $\beta$, both of which are consistent with the $d=\infty$ result (dashed line). Solid lines are guides for the eye.

![Fig. 5.](image)

**Fig. 5.** (A) Dimensional rescaling of the SER (black) and SER breakdown regimes for standard finite-dimensional HS. The early deviation exponent $\omega$ is consistent with hopping in the MK model with $\omega = 0.22$ (red line, Fig. 10), but a growing deviation is observed as $\varphi_c$ increases. (B) The dimensional scaling of HS results for $\varphi_u$, $\varphi_d$, and $\varphi_{\text{SER}}$ converges as $d$ increases, whereas $\varphi_{\text{cset}}$ remains distinctly smaller (compare with Fig. 1C).
Supporting Information

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SI Text

I. Introduction

A. The Model. The infinite-range variant of the Mari–Kurchan (MK) model (1) is defined by adding to the distances between pairs of particles an additional quenched random shift that spans the whole system size (Fig. SI4A). The potential energy contribution to the Hamiltonian for N hard spheres (HS) is thus

\[ \mathcal{H} = \sum_{i<j} U(\|\mathbf{r}_i - \mathbf{r}_j + \mathbf{A}_i\|), \]  

where \( U(r) \) for \( |\mathbf{r}| = r \) is the HS potential \( (e^{-\beta U(r)} = \theta(r - \sigma)) \), for spheres of diameter \( \sigma \), and \( \mathbf{A}_i \) is a uniformly distributed vector within the system volume \( V \), i.e., with a probability distribution \( P(\mathbf{A}_i) = 1/V \). Note that the standard HS model corresponds to \( \mathbf{A}_i = 0 \).

Note that even if in principle all particles interact with all others, in practice because \( U(r) \) is short ranged, a given particle interacts directly only with a finite number of neighbors (the first coordination shell), as in usual liquids. Hence, the model is akin to a mean-field spin-glass model with finite connectivity with the connectivity depending on the number of neighbors in the first coordination shell and thus on both density and dimension.

MK liquids have a simple structure in all spatial dimensions \( d \), because random shifts eliminate higher-order correlations. For example, consider two particles \( i \) and \( j \) both near particle \( k \); i.e., \( |\mathbf{r}_i - \mathbf{r}_j + \mathbf{A}_i| \approx \sigma \) and \( |\mathbf{r}_i - \mathbf{r}_j + \mathbf{A}_j| \approx \sigma \). Unlike in regular HS, in the MK model particles \( i \) and \( j \) have a negligible probability of being near each other \( (|\mathbf{r}_i - \mathbf{r}_j + \mathbf{A}_i| > \sigma) \), because their effective distance is shifted by \( \mathbf{A}_i \), which is of the order of the system size. This argument can also be generalized to interactions between more particles. Particle \( j \) thus has hard-core interactions with its neighbors, but with probability one in the thermodynamic limit these neighbors can overlap with each other. The two-point correlation function seen from one particle is simply

\[ g_2(r) = \left( \frac{1}{N} \sum_{\mathbf{r}_j} \delta(\mathbf{r}_j - r) \right)^N = \theta(\|\mathbf{r} - \sigma\|), \]  

where \( \mathbf{r}_j = \mathbf{r}_i - \mathbf{r}_j + \mathbf{A}_i \).

Let \( V_d(\sigma/2) \) be the volume of a \( d \)-dimensional ball of diameter \( \sigma \), and \( V_d = V_d(1) \). For the MK model, the virial expansion of the equation of state (EOS) terminates at the second-order

\[ p = 1 + B_2 \rho = 1 + 2^{d-1} \varphi, \]

\[ \ln \frac{\rho_{MK}}{\rho_{HS}} = 1 - \log \rho^d - 2^{d-1} \varphi \ln N, \]  

where \( \rho = \beta \rho / \beta \) is the reduced pressure with \( \beta \) the inverse temperature and \( \rho = N/V \) the number density, \( \varphi = \rho V_d(\sigma/2) = \rho V^d/2^d \) is the packing fraction (we set \( \sigma = 1 \)), \( \rho_{MK}^d \) is the liquid entropy per particle, \( \beta \) is the thermal de Broglie wavelength, and \( B_2 = V_d/2 \) is the HS second virial coefficient.

Compared with HS, the MK model has several unique features: (i) Although monodisperse HS easily crystallize in low dimensions, the random shifts in the MK model impose a quenched disorder that is incompatible with crystal symmetry and thus fully suppresses the crystal phase. (ii) Glass–glass nucleation (2, 3) is also suppressed, because if a nucleus forms around a particle, the particles inside this nucleus are actually randomly distributed in real space and thus no surface can be formed (Fig. S1). The free energy cost of forming a nucleus hence scales with the system size and diverges in the thermodynamic limit. (iii) Particle hopping is much less correlated. By contrast to HS, where the hopping of a particle increases the chance that one of its neighbors also hops due to the real-space void it leaves behind, facilitation is limited to unblocking an escape channel in the MK model (Fig. SI6B). (iv) MK particles are distinguishable because the quenched shifts \( \{A_i\} \) are fixed. Besides the lack of structure, the partition function \( Z \) of the MK model is therefore different from that of HS by a factor of \( N! \), i.e., \( Z_{MK}/N! - Z_{HS} \), and hence \( \ln Z_{MK} - \ln Z_{HS} + \ln N \). As a result, the density of the Kauzmann transition in MK diverges in the thermodynamic limit (1).

Introducing a set of quenched random shifts brings two key advantages from a methodological point of view. First, in computer simulations, it is convenient to “plant” an equilibrated MK configuration (SI Text, section I.B.2). Planting avoids the (circular) difficulty encountered in most other glass-forming liquids of equilibrating an initial liquid configuration before studying its equilibrium relaxation dynamics. Second, one can map the model onto a constraint satisfaction problem defined on a random graph (Bethe lattice). It is therefore possible to study its properties with the cavity method (SI Text, section II.A.2), which is, in principle, exactly solvable.

B. Simulation Details. 1. Molecular dynamics simulations. We adapt the event-driven molecular dynamics (MD) algorithm of refs. 4–6 for HS to simulate the MK model in dimensions \( d = 2 - 6 \) with \( N = 4,000 \) particles. Periodic boundary conditions with the minimum image convention are implemented on the shifted distances \( |\mathbf{r}_i - \mathbf{r}_j + \mathbf{A}_i| \). For each \( \varphi \), we perform eight independent realizations, each corresponding to a different set \( \{A_i\} \) for a planted initial configuration (SI Text, section I.B.2). Simulations are run at constant unit \( \beta \), for a time \( t \) (given in units of \( \sqrt{\beta \sigma m^2} \), where the particle mass \( m \) is also set to unity) sufficiently long to reach either the diffusive regime in the liquid or the asymptotic plateau in the glass. As described in refs. 6 and 7, HS data are obtained from simulations of \( N = 8,000 \) identical particles in \( d = 4 - 8 \) and, to prevent the system from crystallizing (8), from a HS binary mixture with diameter ratio \( \sigma_2/\sigma_1 = 1.2 \) in \( d = 3 \) (7).

2. Planting. Planting, which here consists of switching the order of determining initial particle positions \( \{\mathbf{r}_i\} \) and constraints \( \{A_i\} \), is an expedient technique for studying equilibrium ensembles in random constraint satisfaction problems (9). In general, the planted ensemble is different from the annealed ensemble, but for the liquid phase it can be shown that both are equivalent, as we detail below.

In the following, we are interested in physical observables \( \mathcal{F} \) that depend on some initial condition \( \{\mathbf{r}_i\} \) and on their time evolution under deterministic MD dynamics, e.g., the mean square displacement defined in Eq. S7. In the presence of disorder, the average of physical observables should be measured by the so-called “quenched” average

\[ \langle \mathcal{F} \rangle^\Lambda \equiv \int \prod_{i<j}^N \mathcal{A}_i \mathcal{P}(\mathcal{A}_i) \left( \frac{\prod_{i=1}^N d\mathbf{r}_i \mathcal{F} e^{-\beta \mathcal{H}}} {\int \prod_{i=1}^N d\mathbf{r}_i e^{-\beta \mathcal{H}}} \right). \]  

where \( \mathcal{F} \) and \( \mathcal{H} \) depend on both \( \{\mathbf{r}_i\} \) and \( \{A_i\} \). In fact, because the disorder is independent of time for a given sample, one
Because the averages over proportionally to log \( n \). Lately, in the MK model the liquid entropy per particle diverges in the thermodynamic limit. This condition is satisfied in the liquid fluctuations of quenched disorder ages over the disorder are the same, independently, uniformly in the volume \( V \). This describes a situation where both variables and disorder fluctuations are fixed, equilibrating independently, uniformly in the volume \( V \). The last equality in Eq. \( S5 \) shows that the integration measure can be obtained by first extracting a uniformly random configuration \( \{ r_i \} \) and next extracting a configuration \( \{ A_i \} \) from the distribution

\[
P(\{ A_i \} | \{ r_i \} ) = \left[ \prod_{i,j} P( A_i ) \right] e^{- \beta U( |r_i - r_j + A_i| )}.
\]

This average corresponds to a very different situation, where the averages over \( \{ r_i \} \) and \( \{ A_i \} \) are interchangeable. Physically, this describes a situation where both variables and disorder fluctuate together; their timescales are indistinguishable. Mathematically, the last equality in Eq. \( S5 \) shows that the integration measure can be obtained by first extracting a uniformly random configuration \( \{ r_i \} \) and next extracting a configuration \( \{ A_i \} \) from the distribution

\[
P(\{ A_i \} | \{ r_i \} ) = \left[ \prod_{i,j} P( A_i ) \right] e^{- \beta U( |r_i - r_j + A_i| )}.
\]

Because \( P(\{ A_i \} | \{ r_i \} ) \) is factorized, each \( A_i \) must be extracted independently, uniformly in the volume \( V \) with the constraint that \( |r_i - r_j + A_i| > \sigma \). In summary, we use the following procedure to compute \( \langle F \rangle_a \).

**Procedure—planting MK.**

i) Generate \( N \) particle positions \( \{ r_i \} \) according to a Poisson (ideal gas) process.

ii) For each pair of particles \( i \) and \( j \), random and independently draw a vector \( A_{ij} \), uniformly in the subregion of the whole volume \( V \) that is compatible with \( r_i \) and \( r_j \), \( |r_i - r_j + A_{ij}| > \sigma \).

iii) Starting from the state given by \( \{ r_i \} \), and for the given \( \{ A_i \} \), compute the time evolution \( \{ r_i(t) \} \) from MD simulations. From this trajectory compute \( F \).

iv) Repeat steps i–iii to average over disorder and initial configurations.

The key to the success of this approach is determining if, and under what conditions, the quenched and the annealed averages over the disorder are the same, \( \langle F \rangle_a = \langle F \rangle_q \). Eqs. S4 and S5 coincide if the equality \( \log Z_a = \log Z_q \) holds, where \( Z = \int \prod_{i,j} d r_i e^{- \beta H} \) is the partition function for given \( \{ A_i \} \) (9–11). This situation arises if the fluctuations of \( Z \) induced by the fluctuations of quenched disorder \( \{ A_i \} \) are very weak in the thermodynamic limit. This condition is satisfied in the liquid phase, but is violated in the glass phase away from the equilibrium liquid line (9–11).

According to the analysis of ref. 9, to check numerically that the annealed and the quenched average coincide, one should compute the vibrational (internal) entropy of the planted glass state. This can be done, for example, using the procedure described in ref. 12. If the internal entropy of the glass turns out to be larger than the liquid entropy given by Eq. S3, then the annealing average does not coincide with the quenched average (9). Fortunately, in the MK model the liquid entropy per particle diverges proportionally to \( \log N \) (Eq. S3), whereas the glass entropy per particle is finite, because particles cannot exchange (at least if one neglects hopping, as discussed below). Therefore, for \( N \rightarrow \infty \) the liquid entropy is always larger than the glass entropy, and the annealed average is correct. In other words, because the Kauzmann transition for the MK model is located at infinite density (1), the procedure is valid.

Numerical simulations show that the annealed average done using the planting procedure discussed above is in perfect agreement with the liquid EOS, Eq. S3 (Fig. S2). This result is not a surprise, because it can easily be shown that the annealed equation leads to the same liquid EOS in Eq. S3, but it is a consistency test for the numerical procedure. Note that the pressure remains stable over time, as it should be if one initializes the MD simulation in an equilibrium configuration.

**C. Basic Phenomenology of Glassy Behavior and Definitions of Physical Quantities.** Before turning to a more detailed explanation of our results, in this section we summarize the main physical observables that we investigate in this study, with a short account of their definition and of the main results.

1. **Mean square displacement and cage sizes.** Despite its trivial liquid phase, the MK model presents a complex glass-forming and glassy behavior. Above the onset \( \theta_{QUT} \) (SI Text, section I.C.3), we can distinguish three main regimes in the mean square displacement,

\[
\Delta(t) = 1/N \sum_{i=1}^N \langle |r_i(t) - r_i(0)|^2 \rangle.
\]

\( (i) \) a ballistic regime with \( \Delta(t) = dt^2 \); \( (ii) \) a caging regime with a plateau \( \Delta(t) \sim \Delta \), where \( \Delta \) is the mean cage size; and \( (iii) \) a diffusive regime with \( \Delta(t) = 2Dt \), where \( D \) is the diffusivity. According to mode-coupling theory (MCT), the plateau becomes asymptotically stable beyond the dynamical transition \( \theta_d \) (SI Text, section II.B.1). We can then formally define the mean cage size as the infinite time limit of the mean square displacement (MSD),

\[
\Delta \equiv \lim_{t \to \infty} \beta \Delta(t),
\]

and the individual cage size \( \Delta_i \) of each particle \( i \),

\[
\Delta_i \equiv \lim_{t \to \infty} \langle |r_i(t) - r_i(0)|^2 \rangle.
\]

From this definition and the equilibrium conditions \( \langle r_i(0) \rangle = \langle r_i(t) \rangle \) and \( \langle |r_i(0)|^2 \rangle = \langle |r_i(t)|^2 \rangle \), we obtain another expression for \( \Delta_i \),

\[
\Delta_i = \lim_{t \to \infty} \left[ \langle |r_i(t)|^2 \rangle - 2 \langle r_i(t) \cdot r_i(0) \rangle + \langle |r_i(0)|^2 \rangle \right]
\]

\[
= 2 \lim_{t \to \infty} \left[ \langle |r_i(t)|^2 \rangle - |\langle r_i(t) \rangle|^2 \right].
\]

The definition of \( \Delta_i \) in Eq. S9 can be directly used to measure individual cage sizes in numerical simulations. Eq. S10 also suggests that \( \Delta_i \) is twice the variance of the distribution of particle positions within a cage. In theoretical calculations, a cage form ansatz \( f_d(r) \) is usually used for this distribution. Two commonly used functions are the Gaussian

\[
f_d^G(r) = e^{-r^2/(2\Delta)}
\]

and the ball functions

\[
f_d^B(r) = \frac{\theta(A - r^2)}{V_d(\sqrt{A})}
\]
Below, we use the Gaussian ansatz in the replica method (SI Text, section II.A.2) and both ansatzes in the cavity method (SI Text, section II.A.2). The parameter $A$ in these functions can be related to $\Delta_t$, using Eq. S10,

$$\Delta_t = 2d A_t$$

for the Gaussian function and

$$\Delta_t = \frac{2d}{d+2} A_t$$

for the ball function.

2. Characteristic timescales. In this subsection, we define the characteristic timescales, their physical interpretations, and how they are numerically determined.

- $\tau_0$, microscopic time: This natural timescale serves as a reference to compare the evolution of other timescales with spatial dimension $d$. Its definition is such that the velocity autocorrelation function $d(\tau_0) = 1/e$, where $d(t) = (1/dN)\sum_{i=1}^{N} \langle v_i(t) \cdot v_i(0) \rangle = (1/2d)\langle d^2(\Delta(t)/d^2) \rangle$ (Fig. S3C) (13).

- $\tau_D$, diffusion time: The characteristic time for diffusion is defined as $\tau_D = \sigma^2/D$, such that $\Delta(t)$ vs. $t/\tau_D$ collapses in the caging and diffusive regimes (Fig. S3A), as predicted by MCT (Eq. S34). Using this collapse, we can determine $\tau_D$ without explicitly extracting $D$, which allows us to estimate $\tau_D$ close to the dynamical transition, even when the fully diffusive regime itself is beyond numerical reach.

- $\tau_{SER}$, characteristic time at $\phi_{SER}$; i.e., $\tau_{SER} = \tau_D(\phi_{SER})$.

- $\tau_\alpha$, structural relaxation time: In standard glass-forming liquids, $\tau_\alpha$ is typically extracted from the decay of the self-intermediate scattering function

$$F_2(k, t) = \frac{1}{N} \sum_{i=1}^{N} e^{i k \cdot r_i(t) - r_i(0)}$$

such that

$$F_2(k^*, \tau_\alpha) = \frac{1}{e},$$

where $k^*$ is the first particle peak of the structure factor

$$S(k) = 1 + \rho \int dr \, e^{i k \cdot r} g_2(r).$$

For the MK model, however, this method cannot be directly applied because the trivial structure of $g_2(r)$ (and hence of $S(k)$) leaves $k^*$ ill defined. Here, we use a slightly different, although consistent, approach to measuring $\tau_\alpha$. We first generalize the definition of the MSD to the typical displacement of particles

$$r_{typ}(t) = \lim_{z \to 0} \frac{1}{N} \sum_{i=1}^{N} \langle |r_i(t) - r_i(0)|^z \rangle^{1/z},$$

which is the zeroth moment of the self van Hove function $G_v(r, t)$, i.e., the displacement of the majority of particles at time $t$. In practice, to determine $r_{typ}(t)$, we use $z = 0.1$, which is very close to the limit $z \to 0$. By analogy to $\tau_D$, we then extract the relaxation time $\tau_\alpha$ by ensuring that $r_{typ}(t)$ vs. $t/\tau_\alpha$ collapses in the MSD caging regime (Fig. S3B). Note that $\tau_\alpha$ is then defined only up to an overall constant that is independent of density.

For HS, this (re)definition of $\tau_\alpha$ is consistent with the traditional one, because the condition in Eq. S16 is equivalent to $k r_{typ}(\tau_\alpha) \sim 1$. The length scale $1/k^*$ indeed corresponds to that of the maximum density fluctuation, which should be of the order of the typical cage diameter. The scaling $r_{typ}(\tau_\alpha) \sim 1/k^* \sim \sqrt{\tau}$ shows that $r_{typ}$ is near the caging regime at $\tau_\alpha$, and hence should be independent of density. Our estimate of $\tau_\alpha$ is therefore consistent with the proportionality relation for the viscosity $\tau_\alpha \sim \eta$ observed in very sluggish fluids (7). Note that the above definitions of $\tau_\alpha$ and $\tau_D$ give additional weight to slower and faster particles, respectively.

In this context, the breakdown of the Stokes–Einstein relation (SER) is consistent with a proportion of fast particles that is larger than expected (14).

- $\tau_h$, hopping time: The typical time for a caged particle to escape (details in SI Text, section III).

3. Characteristic densities. In this subsection, we define the characteristic densities (number density and volume fraction are used interchangeably), their physical interpretations, and how they are numerically and theoretically determined. Results for HS and the MK model are reported in Table S1.

- $\phi_{onset}$, onset density of the glassy behavior: It corresponds to the lower limit of the caging regime (15). Its choice is such that for $\phi < \phi_{onset}$ no inflection point appears in the logarithm-scale MSD; for $\phi \geq \phi_{onset}$, the MSD shows an inflection point, i.e., a point where $d^2 \ln \Delta(t)/(d \ln t)^2 = 0$. In this regime a non-Fickian behavior is observed. Hence, $\phi_{onset}$ also corresponds to the density at which the minimum value of the non-Fickian coefficient $\nu_{min}$ (Fig. S3D), where $\nu_{min} = \min[\nu(t)]$ and the non-Fickian coefficient $\nu(t) \equiv d \ln \Delta(t)/d \ln t$ (16). Note, however, that our estimate of $\phi_{onset}$ likely underestimates the onset calculated from the emergence of a finite configurational entropy in static calculations (17).

- $\phi_{SER}$, characteristic density for the breakdown of the SER: Below $\phi_{SER}$, hopping is irrelevant because, if present, it is indistinguishable from the regular liquid dynamics, and the MCT scaling relations are satisfied (SI Text, section II.B.1); above $\phi_{SER}$, hopping becomes faster than the characteristic MCT time ($\tau_\alpha < \tau_D$), and consequently both the MCT scaling and the SER are violated.

- $\phi_d$, dynamical glass transition threshold: In the MK model, this density is theoretically calculated from the replica method (SI Text, section II.A.3) and numerically confirmed by testing the MCT scaling $\tau_\alpha \sim |\phi - \phi_d|^{-\gamma}$ (or equivalently, $D \sim |\phi - \phi_d|^\beta$) in the density range over which hopping is negligible ($\phi_{onset} < \phi < \phi_{SER}$). In the HS model, however, we lack reliable theoretical predictions for $\phi_d$ in low dimensions. We therefore determine $\phi_d$ from fitting the simulation results for $D$ in the regime $\phi_{onset} < \phi < \phi_{SER}$. Our results are consistent with those reported in ref. 5, where $\phi_d$ was extrapolated from slowly quenching the fluid. Note that $\phi_d$ is sharply defined only when hopping contributions can be separated without ambiguity. Hence, $\phi_d$ is well defined only in the replica calculation, where hopping is excluded by construction.

- $\phi_{T}$, effective dynamical glass transition threshold: Empirically, $\phi_{T}$ is determined by fitting the diffusivity data, as is commonly done in glass formers. In this study we show, however, that $\phi_{T}$ is systematically shifted with respect to $\phi_d$ ($\phi_{T} > \phi_d$). Note that because $\phi_{T}$ is a fitting parameter, it also depends on the density range one chooses (or that is available) for the power-law fit.

- $\nu_0$, phenomenological parameter from the Vogel–Tammann–Fulcher (VTF) fit to $\tau_D \sim e^{B \nu_0/(\nu_0 - \psi)}$. In the MK model, $\nu_0$ is clearly different from the thermodynamic Kauzmann transition point $\nu_K = \infty$. Recall, however, that the MK model lacks the glass–glass nucleation processes assumed by the Adam–Gibbs (AG) and the random first-order transition (RFOT) theories, to
associate the divergence of the relaxation timescale with the thermodynamic singularity at $q_P$.

- $q_P$: percolation threshold for the cage network: Below $q_P$, a particle can diffuse by successive hops on the percolating network of cages. Because the infinite time limit of the MSD is truly bounded only above this threshold, $q_P$ also provides an upper bound for $q_K$; i.e., $q_K > q_P$.
- $q_K$: Kauzmann transition: density at which the complexity $\Sigma$ (or configurational entropy) vanishes: As discussed above, because $\Sigma_{MK} \sim 2\pi^2 + \ln N$, the density of the Kauzmann transition diverges ($q_K = \infty$) in the thermodynamic limit.

### II. Caging

#### A. Thermodynamics: The Caging Order Parameter and the Dynamic Transition Density

The mean caging order parameter can be obtained equivalently from the replica method, following refs. 18 and 19, or from the cavity method, following ref. 20. Here we briefly describe how these approaches are adapted to the MK model.

1. **Calculation of the mean caging order parameter: the replica method.** References 18 and 19 used the replica approach to obtain HS results, and it is straightforward to check that these derivations rely on the pair correlation function only in the liquid phase; terms corresponding to third- and higher-order structural correlations are neglected. The treatment of ref. 18 can therefore be directly applied to the MK model, for which these assumptions are exact. The results from ref. 19 have also been obtained using the approximation $g_z(\sigma) = y_{0z}(\sigma) \phi(\sigma - \sigma)$ (ref. 19, equation 21) (the soft-sphere temperature is set to zero to study hard-core particles). Comparing this result with Eq. S2, we see that for the MK model $y_{0z}(\sigma) = 1$. All of the results of refs. 18 and 19 can thus be straightforwardly extended to the MK model by setting $y_{0z}(\sigma) = 1$. (Note that the discussion of ref. 19 was restricted to $d = 3$, but a general discussion for all $d$ can be found in ref. 18).

The replicated entropy thus has the form

$$ S(m.A; T, \phi) = S_0(m.A) + \Sigma_{MK}(\phi) + 2^{d-1}qG(m.A), $$

$$ S_0(m.A) = \frac{d}{2}(m - 1)\ln(2\pi A) + \frac{d}{2}(m + 1 + \ln m), $$

$$ G(m.A) = \frac{d}{2} \int_0^\infty dr r^{d-1} |q_A(r)|^m - \theta(r - \sigma), $$

$$ q_A(r) = \int 2\pi f_2^d(r') \theta |r - r'| - \sigma $$

$$ = \int \frac{d\mu}{\sqrt{\pi}} (\sigma - 1/2)^{d-1} e^{-\sigma^2/4\mu} \sqrt{\frac{\sigma^2}{\pi}} I_{d-2}/2 \sqrt{\frac{\pi}{\sqrt{\pi} A}} $$

$$ \times e^{-\pi m/2A} \sqrt{\frac{m}{A}} \langle d-2 \rangle/2 \sqrt{\frac{m}{2A}}, $$

where $f_2^d(r)$ is the $d$-dimensional Gaussian cage given in Eq. S11 and $I_{d-2}$ is the modified Bessel function. The last expression for $q_A(r)$ is obtained using bipolar coordinates to compute the convolution (18). Remarkably, in odd dimensions the integral over $\mu$ can be computed analytically, which facilitates the numerical evaluation of the replicated entropy.

From Eq. S19, we can derive the equation for $A$ from the condition $\partial S/\partial A = 0$, which reads

$$ 1 = \frac{2^d q}{d} \frac{d G(m.A)}{\partial A} = \frac{2^d q}{d} F(m.A). $$

The cage radius in the liquid can be obtained by solving this equation in the limit $m \to 1$, and $\Delta = 2dA$. From Eq. S20, one sees that the dynamical transition $q_P$ corresponds to the point where $(2^d q/d)\max_y F(1.A) = 1$. Note that for $q > q_P$ Eq. S20 admits two solutions, but only the smaller of the two is a stable physical solution (18).

2. **Calculation of the cage size distribution: the cavity method.** More information on the distribution of individual cage shapes and sizes can be obtained from the cavity method (10, 21). Its application to the MK model has been developed in ref. 20, where the cavity equations are derived and discussed. Here, we present only the main steps.

a. **Cavity fields and replica symmetric cavity equations.** In the cavity approach, the system is described by a set of cavity fields $y_j(r)$. Each cavity field describes the probability of finding a particle at position $r$, when it is added to a system of $N - 1$ particles. The replica symmetric cavity equations provide a recurrence equation for determining these cavity fields,

$$ y_j(r_0) = \frac{1}{2z_0} \prod_{j=1}^{N_0} \int 2\pi f_2^d(r) \theta |r - r'| - \sigma d\eta $$

$$ z_0 = \int 2\pi f_2^d(r) \theta |r - r'| - \sigma d\eta $$

In this recurrence, the new particle interacts with the $N_0$ other particles, each described by its own cavity field $y_j(r)$. The interaction is given by the hard-core constraint $\chi(r) = e^{-\beta U(r)} = \theta(r - \sigma)$. In this equation the quenched random variables $\Lambda_0$ are the random shifts that appear in the Hamiltonian, but they should be independently extracted at each cavity iteration. They are independently distributed in the whole volume $V$ with a uniform distribution $P(\Lambda_0) = 1/V$. Note that in ref. 20, the cavity equations were obtained for a model defined on a random graph that is locally tree-like, corresponding to a situation where $N_0$ remains finite as $N \to \infty$. The method, however, is also applicable to the MK model, where $N_0 = N - 1$, corresponding to the fully connected graph (10). A convenient way to obtain the fully connected graph is to first take the limit $N \to \infty$ and then $N_0 \to \infty$. One can show that this procedure is indeed equivalent to considering $N_0 = N - 1$ (10).

b. **Translational invariance and irrelevance of the random shifts.** To describe the liquid and the glassy states of the MK model, we are interested in solutions of the cavity equation that have statistical translational invariance. To be more precise, the liquid phase is described by uniform fields $y(r) = 1/V$ for all particles. Physically, this situation corresponds to particles diffusing everywhere within the system volume, which mathematically reproduces the virial expansion (20). In the glass phase, each individual cavity field has the form $y(r) = f_A(r - R)$, where $f_A(r)$ is a cage function localized around $r = 0$. The cavity field is thus localized around point $R$, but the localization centers $R$ themselves must be uniformly distributed in the whole volume because the glass is globally translationally invariant. Hence, in Eq. S21, when neighbors are picked at random, they are localized around uniformly distributed random positions in space, which makes the random shifts redundant. In the following, we can thus neglect the random shifts and write the replica symmetric cavity equations as

$$ y_0(r_0) = \frac{1}{2z_0} \prod_{j=1}^{N_0} \int 2\pi f_2^d(r) \theta |r - r'| d\eta $$

$$ z_0 = \int 2\pi f_2^d(r) \theta |r - r'| d\eta $$

and take $N_0 \to \infty$. 

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c. The glass phase and the 1RSB cavity equations. In the glass phase, as mentioned above, \( \psi(r) \) are random variables described by a probability distribution \( Q[\psi] \). In the regime that is here of interest, the glass is described by the 1RSB cavity equations derived in refs. 10 and 21. From these equations, we obtain that the probability distribution \( Q[\psi] \) satisfies the self-consistent equation

\[
Q[\psi_0] = \frac{1}{z_0(m)} \int \prod_{j=1}^{N_0} dQ[\psi_j]_0 \delta[\text{Eqs. S22}],
\]

where

\[
Z_0(m) = \int \prod_{j=1}^{N_0} dQ[\psi_j]_0,
\]

which makes explicit that the \( N_0 \) cavity fields describing the neighborhood of the new particle are extracted independently from \( Q[\psi] \). The new cavity field is constructed according to Eq. S22 and weighted according to \( z_0^n \). Note that \( z_0 \) is the free volume associated with the new particle, and therefore, by varying the free parameter \( m \), one can select glassy states according to their free volume or, equivalently, their internal entropy.

d. Reconstruction equations. Beyond the dynamical transition, ergodicity is broken in the liquid phase, which corresponds to the liquid splitting into many distinct glassy states. It is well known, however, that if configurations are sampled with the equilibrium Gibbs–Boltzmann measure, then both entropy and pressure are analytic around \( \psi_0 \) and the equilibrium glass phase is the analytical continuation of the liquid phase. To weight glassy states according to the equilibrium Gibbs–Boltzmann measure, one has to weight them proportionally to their free volume, and hence one must set \( m = 1 \) (21). In the case \( m = 1 \), the 1RSB equations greatly simplify due to a mapping onto the reconstruction formalism (22). Reconstruction is then done by introducing new fields \( R_r[\psi(r)] \equiv \psi(r)Q[\psi(r)] \). This change of variable ensures that only the fields that are localized around point \( r \) contribute to \( R_r[\psi(r)] \). Using the global translational invariance \( R_r[\psi(r)] = R_0[\psi(r - r)] \), we conveniently get

\[
R_0[\psi(r_0)] = \int \prod_{j=1}^{N_0} \left[ \frac{d\psi_j(r_j)}{d\psi_j(r')} \right] dR_0[\psi_j(r_j)] \delta(\ast),
\]

\[
\ast \leftarrow \psi(r_0) - \frac{1}{z_0} \prod_{j=1}^{N_0} \left[ \int d\psi_j(r_j) \psi_j(r_0 - r_j - r_j') \right].
\]

Note that the reweighting term \( z_0^n \) has now disappeared from the equations. Note also that only \( R_0[\psi] \) enters the equations and therefore all cavity fields are localized around the origin. The \( r_j \) in Eq. S24 are random shifts of the cavity fields that are constrained to be outside a sphere of radius \( \sigma \) around the origin. The neighbors \( j \) are thus localized outside that sphere, which guarantees that around the origin there exists a void to accommodate an additional particle.

e. Ansatz on the cage shape. As discussed in ref. 20, numerically solving the cavity equations in Eq. S24 remains a formidable task. Here, we make a simple ansatz on the cage shape to facilitate this computation. We first assume that the cavity fields all have the form \( \psi_j(r) = f_{A_j}(r - R_j) \), where \( f \) is a fixed (spherically symmetric) cage shape. We then choose either a Gaussian (Eq. S11) or a ball (Eq. S12) cage shape, with \( \Delta_0 \) given by Eqs. S13 and S14, respectively. We assume that the cage shapes are distributed according to a function \( P_f(A) \) whereas the centers \( R_j \) are uniformly distributed within the volume, as discussed above. We therefore obtain the ansatz

\[
Q[\psi(r)] = \int dP_f(A) \int dR \delta[\psi(r) - f_A(r - R)],
\]

\[
R_0[\psi(r)] = \psi(0)Q[\psi(r)] = \int dP_f(A) \int dR f_A(R) \delta[\psi(r) - f_A(r - R)].
\]

The above equations show that fields contributing to \( R_0[\psi] \) are localized around a point \( R \) that is distributed according to \( f_A(r) \), and hence \( R \) is itself localized close to the origin. Plugging this ansatz in Eq. S24, we obtain

\[
R_0[\psi(r_0)] = \int \prod_{j=1}^{N_0} \left[ \frac{d\psi_j(r_j)}{d\psi_j(r')} \right] dP_f(A_j) \int dR \delta f_A(R_j) \delta[\psi(r_0 - r_j - R_j)]
\]

where

\[
q_A(r) = \int d\psi f_A(r + \psi) \psi.
\]

Note that the factor of 2 is introduced to follow the notational convention of ref. 18.

f. Reconstruction procedure. The physical interpretation of the reconstruction equation is quite straightforward. To construct a new cavity around the origin, one should draw at random \( N_0 \rightarrow \infty \) particles that are located at random positions \( r_j \) outside a sphere of radius \( \sigma \) around the origin. These particles are themselves within a cage, whose size \( A_j \) is extracted from \( P_f(A) \). The point \( r_0 \) is not the center of the cage, but a point that is typical of the distribution inside the cage. The cage center is therefore at \( r_0 + R \), where the shift \( R \) is extracted from the cage shape \( f_A(R) \). Each neighbor rattles around its cage center at \( r_j + R_j \), producing an effective potential that convolutes the HS constraint with the cage shape, \( e^{-\beta q_A(r)} = q_{A_j}(r_0 - r_j + R_j) \). The new cavity field is then given by the (normalized) exponential of the sum of all effective potentials, \( \psi(r_0) \propto \exp[\sum q_{A_j}(r_0 - r_j + R_j)] \). Finally, we note that although the number of neighbors \( N_0 \) should be sent to infinity, distant neighbors do not affect the new cavity field because \( q_{A_j}(r) \) tends to 1 when \( r \rightarrow \infty \). We can therefore introduce an arbitrary spatial cutoff and consider only the neighbors (whose number distribution is Poissonian) that are within this cutoff and then increase the cutoff until the results converge. This approach is expressed by the following recursive procedure for self-consistently determining the distribution \( P_f(A) \), which is the only remaining unknown in the cavity reconstruction. Note that once \( P_f(A) \) has been calculated, one can easily obtain the distribution of mean square displacements in the cage, \( P_f(\Delta) \), according to Eq. S13 or Eq. S14. This observable is also easily measured in numerical simulations (and experiments).

Procedure—reconstruction MK.

i) Consider a spherical shell \( \sigma < r < \sigma + \sigma_{cut} \) of volume \( V_0 \) (the upper bound \( \sigma_{cut} \) should be sufficiently large for the results to be independent of it). Consider a number of centers \( N_0 \) distributed according to a Poisson law with average \( N_0 = \nu V_0 \). Uniformly draw these sphere centers \( r_i \) within the shell.

ii) Independently draw \( N_0 \) cage radii \( A_j \) from \( P_f(A) \) and \( N_0 \) displacements \( R_j \) from \( f_A(R_j) \).

iii) From these \( N_0 \) random variables, derive a new cavity field

\[
Q[\psi(r)] = \int dP_f(A) \int dR \delta[\psi(r) - f_A(r - R)],
\]

\[
R_0[\psi(r)] = \psi(0)Q[\psi(r)] = \int dP_f(A) \int dR f_A(R) \delta[\psi(r) - f_A(r - R)].
\]

The above equations show that fields contributing to \( R_0[\psi] \) are localized around a point \( R \) that is distributed according to \( f_A(r) \), and hence \( R \) is itself localized close to the origin. Plugging this ansatz in Eq. S24, we obtain

\[
R_0[\psi(r_0)] = \int \prod_{j=1}^{N_0} \left[ \frac{d\psi_j(r_j)}{d\psi_j(r')} \right] dP_f(A_j) \int dR \delta f_A(R_j) \delta[\psi(r_0 - r_j - R_j)]
\]

where

\[
q_A(r) = \int d\psi f_A(r + \psi) \psi.
\]
tion III.A. We include then check whether any hopping has occurred during this trajectory. Once hopping is removed, our results indicate that the particle is integrated into the effective potential and their vibrational contribution to the motion of the caged cavity parameter is chosen to construct a new cavity solution properly converges when \( \varphi > \varphi_d \) (Fig. S4). We represent the distribution \( P_f(A) \) by a number \( N = 10^6 \gg N_0 \) of samples \( A_k, k = 1 \cdots N \).

3. Comparing theoretical predictions with simulations. We first show that the mean square displacement \( \bar{\Delta} \) predicted from both the replica and the cavity methods is generally in good agreement with the simulation data (Fig. S5). The simulation \( \bar{\Delta} \) is extracted from the asymptotic time limit of the MSD data, according to the MCT scaling (Eq. S39). Close to \( \varphi_d \), the replica theory predicts a scaling (Fig. S5)

\[
|\bar{\Delta}(\varphi) - \Delta_d| \sim |\varphi - \varphi_d|^{1/2}.
\]

where \( \Delta_d = \bar{\Delta}(\varphi_d) \), which is consistent with the MCT prediction. However, around \( \varphi_d \) precisely determining \( \bar{\Delta} \) from either simulations or cavity reconstruction requires a careful consideration of hopping. The simulation and the cavity data therefore unsurprisingly deviate from Eq. S33 in that regime (Fig. S5).

Cavity reconstruction provides a theoretical prediction for the distribution \( P_f(\Delta) \) of individual mean square displacements. To obtain individual cages from simulation, we use Eq. S10 at \( t = 2 \), which is sufficiently long for the cages to form, but not so long that a large fraction of particles have hopped. Note that at densities well above \( \varphi_d \), hopping is so rare that this choice of timescale is irrelevant. As discussed in the main text, our theoretical results agree well with simulations and are independent of the Gaussian or of the ball ansatz for the shape functional. The replica calculation for \( \bar{\Delta} \) using the Gaussian functions and \( P_f(\Delta) = \delta(\Delta - \bar{\Delta}) \) also agrees with the simulation results (Fig. S5).

In summary, we find a basic consistency between our MD simulations and theoretical calculations, including (i) the replica calculation with a Gaussian ansatz for the cage shape and a \( \delta \)-function approximation for the cage size distribution function \( P_f(\Delta) \approx \delta(\Delta - \bar{\Delta}) \) and (ii) the cavity method with both Gaussian and ball ansatzes. It has been shown that in the limit \( d \to \infty \), the theoretical result (of replica calculation) is independent of the cage shape ansatzes \( (23) \), and we also expect it to be independent of the method we use (replica/cavity). In finite dimensions, a weak dependence is expected, but according to our results presented here, it is insignificant compared with the numerical accuracy of the resolution of the cavity equations.

8. Caging Dynamics: MCT and Beyond. In this section, we compare the MD results with the dynamical caging behavior predicted by MCT. The MCT scalings are found to be consistent with our data only when \( \varphi < \varphi_{\text{ser}} \). Above \( \varphi_{\text{ser}} \), MCT predictions are violated, which is well captured by the breakdown of SER and is a consequence of entangling cavity hopping, as discussed in SI Text, section III. It is important to note that we here refer to MCT only as the general scaling laws predicted by the schematic MCT equation \( (24) \), which can be also independently derived from the static framework \( (25) \). The traditional MCT kernel being incorrect for the MK model \( (1) \), the numerical MCT predictions are indeed unsuitable for comparison.

1. Testing the mode-coupling theory. We first compile the MCT predictions tested in our study. The derivations of these predictions as well as many important physical interpretations can be found in ref. 24 and references therein. We denote \( e = (\varphi - \varphi_d)/\varphi_d \) as the distance from the dynamical transition and \( \tau_\beta \) as the characteristic time for the \( \beta \) relaxation. Note that MCT does not predict any breakdown of the SER, so we do not distinguish between the \( \alpha \)-relaxation time \( \tau_\alpha \) and the diffusion time \( \tau_D \) in this analysis \( (\tau_D \approx \tau_\alpha \gg \tau_\alpha) \) Below the dynamical transition \( \varphi < \varphi_d \), MCT predicts that the time evolution of the MSD has the form...
Eq. \( \Delta(t) = \begin{cases} \Delta_d - B|t|^{1/2} \left( \frac{1}{\tau_e} \right)^a, & t \ll \tau_e, \\ \Delta_d + C \left( \frac{t}{\tau_d} \right)^b + \frac{1}{\tau_d}, & t \gg \tau_e, \end{cases} \) \[S34\]

where \( B \) and \( C \) are density-independent constants, and the exponents \( a \) and \( b \) are related by the exponent parameter \( \lambda \) as

\[ \lambda = \frac{\Gamma(1-a)^2}{\Gamma(1-2a)} = \frac{\Gamma(1+b)^2}{\Gamma(1+2b)} \] \[S35\]

Eq. \( S34 \) shows that the relaxation process of \( \Delta \) can be divided into three regimes: \( i \) an early \( \beta \)-relaxation toward the plateau \( \Delta_d, \Delta_d - \Delta_\alpha(t) \sim t^{-\alpha}; \) \( ii \) a late \( \beta \) relaxation leaving from the plateau, \( \Delta_\alpha(t) \sim t^{1/\lambda}; \) and \( iii \) a diffusive process that is linear in time, \( \Delta(t) \sim t. \) We stress that, as described in Section I.C.1, before regime \( i \), there is a ballistic regime characterized by a microscopic time that is much smaller than \( \tau_e \) and is not included in Eq. \( S34 \).

One of the most important predictions made by MCT is that, upon approaching \( \varphi_d \), a power-law divergence should be observed for

\[ \tau_D \sim |\varphi - \varphi_d|^{-\gamma} \] \[S36\]

and

\[ \tau_e \sim |\varphi - \varphi_d|^{-1/2a} \] \[S37\]

where the exponents are related via

\[ \gamma = \frac{1}{2a} + \frac{1}{2\lambda} \] \[S38\]

Beyond \( \varphi_d \), MCT then predicts

\[ \Delta_\alpha(t) = \begin{cases} \Delta_d - B|t|^{1/2} \left( \frac{1}{\tau_e} \right)^a, & t \ll \tau_e, \\ \Delta_d, & t \gg \tau_e. \end{cases} \] \[S39\]

These scalings are tested by the following procedure.

**Procedure—testing MCT.**

\( i \) Obtain \( \varphi_d \) and \( \Delta_d \) from the replica calculation.

\( ii \) Fit \( \tau_d \) according to Eq. \( S36 \) (Fig. 1 of main text) with the theoretical \( \varphi_d \) to obtain the exponent \( \gamma \). A consistent power-law scaling is observed only below some density \( \varphi_{SER} \); above \( \varphi_{SER} \), \( \tau_d \) becomes smaller than the MCT predictions, implying that an additional relaxation process starts to interfere with the dynamics. This observation suggests that when we fit the diffusivity data, only the data below \( \varphi_{SER} \) should be used. If instead we treat \( \varphi_d \) as a fitting parameter for the entire density range, then we end up with shifted values \( \tilde{\varphi}_d \) and \( \tilde{\gamma} \),

\[ \tau_D \sim |\varphi - \tilde{\varphi}_d|^{-\tilde{\gamma}}. \] \[S40\]

From the analysis presented in the main text, it is clear that MCT actually fails when \( \varphi > \varphi_{SER} \), and thus the apparent power-law fitting of Eq. \( S40 \) is not reliable. Results for \( \varphi_d \), \( \tilde{\varphi}_d \), \( \gamma \), and \( \tilde{\gamma} \) can be found in Table S1.

\( iii \) Determine \( a \) and \( b \) (Table S1) from \( \gamma \), using Eqs. \( S35 \) and \( S38 \).

\( iv \) Test the dynamical behavior of \( \Delta(t) \) (Eq. \( S34 \)) below \( \varphi_d \) and determine the constants \( B \) and \( C \). Note that here we have fixed all of the other parameters, \( \varphi_d, \Delta_d, a, \) and \( b \) from previous steps. Fig. S6 \( A \) and \( B \) shows that when \( \varphi < \varphi_{SER} \), Eq. \( S34 \) is satisfied in the entire time regime; when \( \varphi < \varphi_{SER} \), it is satisfied only in the early \( \beta \)-relaxation regime.

\( v \) Using the same constant \( B \), check the MCT dynamics above \( \varphi_d \), using Eq. \( S39 \). When \( \varphi \) is not too far away from \( \varphi_d \), \( \Delta(t) \) does not strictly saturate to a plateau as predicted by MCT, and the scaling behavior of the intermediate time regime is modified.

2. **Breakdown of the Stokes–Einstein relation.** The above analysis shows that the MCT scalings start to break down close to the dynamical transition \( \varphi_d \). To further investigate this property, we look at the scaling relation between the diffusion time \( \tau_d \) and the relaxation time \( \tau_e \). If SER were obeyed, we should obtain \( \tau_D \sim \tau_e \). As shown in Fig. 1 in the main text, SER breaks down when \( \varphi > \varphi_{SER} \) as

\[ \tau_D \sim t^{1-\omega}. \] \[S41\]

where the exponent \( \omega = 0.22 \) is invariant with \( d \).

The breakdown of SER is beyond the MCT description. Interestingly, we observe that three phenomena happen at the same density \( \varphi_{SER} \); \( i \) violation of MCT scalings, \( ii \) violation of SER, and \( iii \) the hopping characteristic time \( \tau_h \) becoming comparable with \( \tau_D \). Our interpretation of these observations is presented in the main text.

C. **Percolation of the Cage Network.** Because cages can be connected via hopping channels, it is natural to examine how the cages are topologically connected. We find that the network of cages spans the system below density \( \varphi_p \). Above \( \varphi_p \), only local cage clusters are formed and particles become strictly confined. We show that this phenomenon can be mapped onto a void percolation transition, which belongs to the same universality class as regular percolation.

1. **Mapping the glass transition to a void percolation transition.** To do the mapping, we first consider the simplest case, where we assume that all of the neighbors of a given particle are frozen, \( P_f(A) = \delta(A) \). We want to know whether the caged particle can move to another cage without overlapping with other particles. Equivalently, we can rescale the size of neighbors as \( \sigma \rightarrow 2\sigma \) and look for a hopping path for the point representing the caged particle in the leftover void space (Fig. S7A).

We next consider the situation where cage sizes are not zero. In this case, particle \( f \) is rattling inside a cage with radius \( \sqrt{A_}\), whose distribution is a density-dependent function \( P_f(A) = \delta(A) \). If a certain channel were closed in the first case, there is now a possibility for it to be open because the particles bounding that channel are now thermally moving. Because we are interested in the upper bound for percolation, i.e., the best-case scenario for hopping, we rescale particle sizes as (Fig. S7A)

\[ \sigma \rightarrow 2(\sigma - 2\sqrt{A}) \] \[S42\]

where \( A \) is drawn from \( P_f(A) \). If no path in void space is found by this construction, then the particle is confined. Strictly speaking, this procedure works only for cage shapes with sharp boundaries, like the ball function in Eq. S12. For a Gaussian cage in Eq. S11, the confined particle always has a finite (but vanishingly small) probability to hop, even if the cage is found to be closed in the percolation mapping. In the following percolation analysis, to avoid any possible confusion, we assume that all cages have ball shapes, which corresponds to assuming that this probability tail is negligible.
2. Methodology for determining the void percolation threshold.

Mapping the void space onto a network via the Voronoi tessellation.

For void percolation, unlike for lattice percolation, or for continuous percolation (which is the dual problem to void percolation), the predefined network is not trivial to extract. It has been shown, however, that the void space between monodisperse spheres can be represented by a network obtained by Voronoi tessellation (26). This method can also be generalized to polydisperse systems via digital Voronoi tessellation (27). In the network representation, nodes are Voronoi vertices, and links are the edges of the Voronoi polyhedra. If any link passes through one or more spheres, then it is blocked and should be removed from the network. After this network is constructed, we check whether there exists a percolated path from the center of the network to the system boundary.

Algorithm for radial Voronoi tessellation of polydisperse spheres in any d. Because our systems have a range of cage sizes, i.e., they map onto spheres with polydisperse diameters, we develop a method to produce the radial Voronoi tessellation for a given configuration. The basic idea is to map the radial Voronoi tessellation in dimension d to a Voronoi tessellation in dimension d + 1 and then use Ohull (28) to compute the Voronoi tessellation.

For the standard Voronoi tessellation, the Voronoi cell for sphere i consists of space points r that satisfy the relation

$$|r - r_i| < |r - r_j|,$$

for any j ≠ i. The radial Voronoi tessellation is a generalization of this definition for unequal-sized spheres,

$$|r - r_i^2| - R_i^2 < |r - r_j^2| - R_j^2,$$

where $R = \sigma/2$ is the particle radius.

To map the radial Voronoi tessellation to a Voronoi tessellation, we denote $R_{\text{max}}$ the maximum radius and introduce a set of points in dimension $d + 1$, $\tilde{r}_i = \left( r_i^1, r_i^2, \ldots, r_i^d, \sqrt{R_{\text{max}}^2 - R_i^2} \right)$, where $i = 1, \ldots, N$. The first $d$ coordinates of $\tilde{r}_i$ are the same as $r_i$, and the final coordinate is a function of the sphere radius. We further introduce a set of dual points $\tilde{r}_i' = \left( r_i^1, r_i^2, \ldots, r_i^d, -\sqrt{R_{\text{max}}^2 - R_i^2} \right)$, as images of $\tilde{r}_i$ with respect to the last coordinates. For each pair $\{\tilde{r}_i, \tilde{r}_i'\}$, we find the $d$-dimensional polygon $P_i$ that is the common boundary between these two points (Fig. S7B). According to the definition in Eq. S43, it is clear that any point $\tilde{r}$ in $P_i$ can be written as $\tilde{r} = (r, 0)$, and $\tilde{r}$ satisfies

$$|\tilde{r} - \tilde{r}_i| < |\tilde{r} - \tilde{r}_i'|,$$

which is equivalent to

$$|r - r_i| + R_{\text{max}}^2 - R_i^2 < |r - r_j| + R_{\text{max}}^2 - R_j^2.$$

Because this relation is exactly the definition of the radial Voronoi cell in Eq. S44, we have proved that the $d$-dimensional polygon $P_i$ is the radial Voronoi cell of sphere $i$ in the original configuration.

Determining the percolation threshold from the scaling theory. The percolation threshold can be determined by finite-size scaling (29). Note that in the void percolation analysis, the variable of interest is the volume fraction of void space $\eta$ and not directly the volume fraction $\phi$. Because our planted configuration is essentially a Poisson process of overlapping spheres, however, we have

$$\eta(\phi) = \left[1 - \frac{V_d(\sigma/2)}{V}\right]^N \approx e^{-\left[N\eta(\sigma/2)/V\right]} = e^{-\phi}.$$

Let $\eta_p = e^{-\psi}$ be the percolation threshold in the infinite systemsize limit. For a system of finite linear size $L \sim V^1/d$, the average effective percolation threshold $\eta_p(L)$ and its variance $\Delta\eta_p(L)$ are linearly related (31),

$$|\eta_p(L) - \eta_p| \sim \Delta\eta_p(L),$$

which allows one to numerically determine $\eta_p$. Because the polydispersity associated with the cage distribution $P_j(A)$ varies with density $\phi$, we cannot, however, directly use this relation. At each $\phi$, we instead modify the rescaling factor of Eq. S42 by adding a factor $K(\phi, L)$,

$$\sigma \rightarrow 2(\sigma - 2\sqrt{A_p})K(\phi, L).$$

and use a binary search to find the percolation threshold $K_p(\phi, L)$ for each configuration. We then calculate $K_p(\phi, L)$ and $\Delta K_p(\phi, L)$ over 1,000 independent realizations and use a similar relation

$$|K_p(\phi, L) - K_p(\phi)| \sim \Delta K_p(\phi, L),$$

to determine $K_p(\phi)$ (Fig. S7C). We finally compute $\eta_p$, such that $K_p(\eta_p) = 1$. Our system is thus percolated at $\eta_p$, without the extra rescaling factor $K_p(\phi)$ (Fig. S7C).

To check the universality of the percolation transition, we examine the scaling of the mean cluster size $\langle V_{\text{net}} \rangle$, where $V_{\text{net}}$ is the total volume of the cluster of cages connected to the planted central cage. According to percolation theory, $\langle V_{\text{net}} \rangle$ diverges at the percolation threshold as a power law with exponent $\gamma$:

$$\langle V_{\text{net}} \rangle \sim |\eta(\phi) - \eta_p|^{-\gamma}. $$

Fig. S7D shows that our results are in agreement with $\gamma = 1.8$ (31) given by lattice percolation, in support of the two problems sharing a same universality class. Based on the above discussion, we summarize the procedure for determining $\eta_p$.

Procedure—$\eta_p$ determination.

i) For a given density $\phi$, obtain the distribution $P_j(A)$ from the cavity method.

ii) Plant a configuration with linear system size $L$ such that the central particle is compatible with all neighbors [this requirement is the same as for the cavity method (SI Text, section II.A.2)].

iii) Rescale the particle sizes following Eq. S49.

iv) Find the percolation threshold $K_p(\phi, L)$ for the configuration, using a binary search. To determine if the void space is percolated:

- Map the $d$-dimensional configuration of rescaled polydisperse spheres to a $(d + 1)$-dimensional configuration of monodisperse spheres.
- Calculate the Voronoi tessellation of the $(d + 1)$-dimensional percolation configuration.
- Map back the $(d + 1)$-dimensional Voronoi tessellation to the $d$-dimensional radial Voronoi tessellation.
- From the radial Voronoi tessellation, construct a network.
- Determine whether the network percolates.

v) Repeat steps i–iv to get $K_p(\phi, L)$ and $\Delta K_p(\phi, L)$.

vi) Vary $L$ and repeat step v to get $K_p(\phi, L)$ and $\Delta K_p(\phi, L)$ at different $L$, and use the finite scaling Eq. S50 to obtain $K_p(\phi)$.

vii) Vary $\phi$ and repeat step vi to get $K_p(\phi)$ at different $\phi$, and find $\eta_p$ such at $K_p(\phi_p) = 1$. 
III. Hopping

In this section, we detail how we detect hopping events in numerical simulations and describe the hopping dynamics of the MK model at a phenomenological level. Theoretical investigations are left for future study.

A. Detecting Hopping. We follow the algorithm of refs. 32 and 33 to detect hopping events in both the MD simulations and the numerical evaluation of the cavity equations. Below we briefly summarize the procedure.

Procedure—detection of hopping.

i) Run simulations and save particle trajectories.

ii) Determine the cage size $\Delta_i$ of each particle as discussed in SI Text, section I.C.1. (We relax the assumption of refs. 32 and 33 that all cages have the same size.)

iii) Split each single-particle trajectory $X(0 < t < t_{\text{tot}})$ in two subsets $X_i(0 < t_i < t')$ and $X_i(t' < t_i < t_{\text{tot}})$ and measure the mean square distance between the two subtrajectories

$$\delta^2(t) = x^2(t_i) \left( d_1^2(t_i) + d_2^2(t_i) \right)^{1/2}, \quad [S52]$$

where $d_i(t_i)$ is the distance between the point at time $t_i$ and the center of mass of the subset $X_i(j, k = 1, 2)$, and $x(t_i) = \sqrt{t_{\text{tot}}-t_i}$ is a normalization factor. Find the time $t^*_{\text{max}}$ such that $\delta^2(t^*_{\text{max}})$ is maximum.

iv) For particle $i$, if $\delta^2(t^*_{\text{max}}) > \Delta_i$, hopping is detected, and the process is repeated recursively for each subtrajectory until $\delta^2(t^*_{\text{max}}) < \Delta_i$ in each subtrajectory.

Following this procedure, we save a sequence of hopping times. Note that in this study we are interested only in the time of the first hopping, which is equivalent to the time during which the particle is trapped in a cage before escaping. Because facilitation is reduced in the MK model, especially at high $\varphi$, we do not specifically distinguish between the first and the subsequent hopping events. The algorithm generally works well at densities $\varphi > \varphi_0$, as shown in Fig. S8. Close to $\varphi_0$, however, hopping is mixed with other relaxation processes, and cages are not clearly defined. Detecting hopping indeed then becomes more sensitive to the specific cutoff thresholds.

B. Hopping Dynamics. Empirically, we find that the above detected hopping time $\tau$ follows a power-law distribution (Fig. 3 of the main text)

$$p_\text{h}(t) \sim t^{-\mu}, \quad [S53]$$

with exponent $\mu < 1$. We can write its cumulative distribution function as

$$G_{\text{h}}(t) = \left( \frac{t}{\tau_h} \right)^{1-\mu}, \quad [S54]$$

where $\tau_h$ is the characteristic hopping timescale, representing the time needed for all particles to hop, $G_{\text{h}}(\tau_h) = 1$. As shown in Fig. 3 of the main text, both $\mu$ and $\tau_h$ depend on $\varphi$. In particular, $\tau_h$ is roughly an exponential function of $\varphi$,

$$\tau_h \sim e^{\varphi_0}, \quad [S55]$$

which suggests that there is no diverging density for $\tau_h$.

Fig. S1. (A) Illustration of the MK model. (Left) Particles in the original space. (Right) Particles in the shifted space with respect to particle i (red). Although neighbors cannot overlap with particle i, they are allowed to overlap with each other because of the random shifts. (B) Comparing hopping in the HS and the MK models. (Left) In HS, removal (hopping) of a neighbor (blue) creates an open channel for the caged particle (red) to hop. (Right) In MK, removing a neighbor is much less likely to open a channel because the other neighbors are allowed to overlap.

Fig. S2. Comparison between the reduced pressure $p$ of planted states (red circles) and the liquid EOS Eq. 3 (black solid line) in $d = 3$. The regime above $\phi_d = 1.776$ is numerically inaccessible from conventional slow quenching procedures [see, for example, the green line obtained from the Lubachevsky–Stillinger algorithm with a growth rate $\gamma = 3 \times 10^{-5}$ (5)], because the system starts to deviate from the liquid EOS around $\phi_d$. (Inset) Planting at $\phi = 1.78$ gives the correct equilibrium pressure (red line) from $t = 0$. 

References
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Fig. S3.  (A and B) Rescaled plots of (A) the MSD and (B) the typical displacement $r^2_{\text{typ}}(t)$ in $d=3$. From right to left, $\varphi = 0.40, 0.60, 1.00, 1.30, 1.40, 1.50, 1.60, 1.65, 1.70, 1.72, 1.74, 1.76, 1.78, 1.80, 1.82,$ and $1.84$, with $\tau_\alpha$ normalized such that $\tau_\alpha \sim \tau_D$ at $\varphi_{\text{onset}}$. Note that the rightmost line (red), at $\varphi_{\text{onset}}$, does not exhibit any plateau regime. (C) Velocity autocorrelation function $d(t)$ at $\varphi_{\text{onset}}$ in $d=2-6$. By definition, $d(\tau_0) = 1/e$ (dashed line). (D) Minimum value of the non-Fickian coefficient $\nu_{\text{min}}$ in $d=3$. At high densities the $\nu_{\text{min}}$ decreases linearly with $\varphi$ (solid line), and at low densities $\nu_{\text{min}} = 1$ (dashed line). The crossover occurs around $\varphi_{\text{onset}} = 0.40(5)$.

Fig. S4. Evolution of mean square displacement $\Delta$ under iteration of the cavity reconstruction, at (solid lines, from bottom to top) $\varphi = 2.50, 2.20, 1.95, 1.80, 1.75, 1.70,$ and $1.60$ in $d=3$. The solution becomes completely unstable above $\Delta_\nu = 0.267$ (dotted black line), as predicted by the replica method. If hopping is not removed, the solution diverges quickly when $\varphi$ approaches $\varphi_d$. See, for instance, the unfiltered results for $\varphi = 1.95$ (pink dashed line).
Fig. S5. (A) The $d = 3$ mean square displacement $\Delta_4(\varphi)$ obtained from the replica method (black line), cavity reconstruction (blue squares), and MD simulations (red crosses). The replica result for $\varphi_d$ corresponds to the point where the theoretical replica line has a square-root singularity. (B) The theoretical results are consistent with the scaling form in Eq. S33, but deviations are observed in the MD data close to $\varphi_d$, due to the ambiguity in determining cage sizes when hopping is significant.

Fig. S6. Testing MCT scalings for the MK model in $d = 6$. (A and B) Below $\varphi_d$, the MD data for $\varphi = 0.30, 0.35, 0.40, 0.41, 0.42, 0.43, 0.435,$ and $0.44$ are fitted to Eq. S34 (red lines) for (A) the early $\beta$ relaxation, and (B) the late $\beta$ relaxation together with diffusion, using fitting parameters $B = 0.073$ and $C = 1.3$. (C) Above $\varphi_d$, the MD data $\varphi = 0.45, 0.455, 0.46,$ and $0.47$ are compared with the early $\beta$-relaxation scaling in Eq. S39 with the same value of $B$. A good agreement is found for the entire time regime when $\varphi < \varphi_{SER}$ (black solid lines). When $\varphi > \varphi_{SER}$ (green dashed lines), we observe a good agreement only for the early $\beta$-relaxation regime, which suggests that at later times hopping mixes with the MCT dynamics.
Fig. S7. Percolation analysis. (A) Mapping the glass transition problem to a void percolation transition according to the rescaling law $\sigma \rightarrow 2\sigma$ (from Left to Center) and according to $\sigma \rightarrow 2(\sigma - 2\sqrt{A_j})$ (from Left to Right). In the latter case, additional hopping paths may be found. (B) Calculating the radical Voronoi cell $P_i$ (blue triangle) of sphere $i$ in dimension $d$ ($d = 2$ in this example). The problem is mapped onto a Voronoi tessellation in dimension $d + 1$, where the tetrahedron is the Voronoi cell of $i$ in the mapped $d + 1$ configuration. (C) Determining $\phi_p$ in $d = 3$. (Left) Estimation of the percolation threshold $K_p(\phi)$ from the finite-size analysis of Eq. S50. (Right) Determining the percolation density $\phi_p = 2.4$ from $K_p(\phi_p) = 1$. The black line is an exponential fit of the data points. (D) Scaling of the mean cluster volume is consistent with Eq. S51, using the exponent $\gamma_p = 1.8$ given by standard lattice percolation.

Fig. S8. An example of hopping detection in $d = 2$ at $\varphi = 2.40$. (A) The particle trajectory clearly reveals two well-formed cages. (B) The hopping between cages is visualized in the time series, with the two detected hopping times (dotted lines) at $t = 3,948.0$ and $t = 7,863.6$. 

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Table S1. Numerical values of characteristic densities and MCT exponents for the MK and the HS models

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<th>$\bar{\phi}_d$</th>
<th>$\phi_0$</th>
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Data for the MK model in $d = 2$–6 and for the HS model in $d = 3$–8. Theoretical results are reported for $\phi_d$ of the MK model, but all of the other values are from simulations. In $d = 8$ for the HS model, no SER violation is detected in the dynamical regime that is computationally accessible.