On the jamming phase diagram for frictionless hard-sphere packings

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We computer-generated monodisperse and polydisperse frictionless hard-sphere packings of $10^4$ particles with log-normal particle diameter distributions in a wide range of packing densities $\varphi$ (for monodisperse packings $\varphi = 0.46-0.72$). We equilibrated these packings and searched for their inherent structures, which for hard spheres we refer to as closest jammed configurations. We found that the closest jamming densities $\varphi_1$ for equilibrated packings with initial densities $\varphi \leq 0.52$ are located near the random close pack limit $\varphi_{RCP}$: the available phase space is dominated by basins of attraction that we associate with liquid. $\varphi_{RCP}$ depends on the polydispersity and is $\sim 0.64$ for monodisperse packings. For $\varphi > 0.52$, $\varphi_2$ increases with $\varphi$: the available phase space is dominated by basins of attraction that we associate with glass. When $\varphi$ reaches the ideal glass transition density $\varphi_g$, $\varphi_3$ reaches the ideal glass density (the glass close pack limit) $\varphi_{GCP}$, so that the available phase space is dominated at $\varphi_g$ by the basin of attraction of the ideal glass. For packings with sphere diameter standard deviation $\sigma = 0.1$, $\varphi_{GCP} \approx 0.655$ and $\varphi_g \approx 0.59$. For monodisperse and slightly polydisperse packings, crystallization is superimposed on these processes: it starts at the melting transition density $\varphi_m$ and ends at the crystallization offset density $\varphi_{off}$. For monodisperse packings, $\varphi_m = 0.54$ and $\varphi_{off} = 0.61$. We verified that the results for polydisperse packings are independent of the generation protocol for $\varphi \leq \varphi_g$.

1. Introduction

Frictionless hard-sphere packings represent a useful model for atomic systems, liquids, glasses, and crystals,1 aside from being a system directly utilized in materials science and chemical engineering.2,3 This simple yet powerful model exhibits a range of diverse phenomena, including melting and freezing transitions,1,4-9 the ideal glass transition,1,7,10-11 the ideal glass or the glass close packing (GCP) limit,1,14 as well as the random close packing (RCP) limit.1,14-16

There are several attempts to merge the multitude of these effects into a single picture.1,18 It is a difficult task, as significant debate on some of the concepts above is underway. The first big challenge is the definition and determination of the RCP limit. For monodisperse particles, there exist at least three estimates for the RCP limit, with distinct densities $\varphi$: (i) $\varphi = 0.634-0.636$,1,16-18 (ii) $\varphi = 0.64$15,19-21 and (iii) $\varphi = 0.65.22-29$ In our previous studies,14,19 we suggested that $\varphi = 0.64$ and $\varphi = 0.65$ refer to different phenomena and represent the RCP limit $\varphi_{RCP}$ (in the sense of the J-point19) and a lower bound of the GCP limit $\varphi_{GCP}$, respectively. It implies that random jammed packings can systematically be produced at any density in the range [$\varphi_{RCP}, \varphi_{GCP}$].1,19,21 The definition and determination of the GCP limit and the corresponding ideal glass transition represent the second actively discussed topic,1,10,13 stemming from research on glasses and colloids. It is debated whether the ideal glass exists and if so, what is the density of the ideal glass transition. It is unclear if there are multiple glassy states and what is the lowest glass transition density.

We believe that the RCP and GCP limits shall be studied together, as a part of the systematic investigation of the phase space structure for hard spheres. At each packing density $\varphi$, the phase space has areas corresponding to valid packing configurations. These areas comprise basins of attractions of jammed configurations.31-35 Some of these basins dominate the available phase space. Thus, one of the characteristics of the phase space at a given $\varphi$ is the jamming density of these dominant basins of attraction, $\varphi_J$. The main objective of this paper is to build a map from $\varphi$ to $\varphi_J$ for a wide range of initial densities $\varphi$.

With this intention to study the structure of the phase space for hard spheres from first principles, we computer-generated monodisperse and polydisperse frictionless hard-sphere packings of $10^4$ particles (cf. Fig. 1) over a wide range of densities $\varphi$ (for monodisperse packings $\varphi = 0.46-0.72$). Polydisperse particles have log-normal diameter distribution with diameter standard deviation $\sigma$ from 0.05 to 0.3 in steps of 0.05. Then, we equilibrated these packings to let packing configurations arrive at the basins of attraction of inherent structures31-35 that dominate the phase space at given densities. Finally, we searched for the inherent structures of these equilibrated...
configurations. In this paper, we use for hard spheres the term “closest jammed configurations” instead of “inherent structures”. An inherent structure for an arbitrary configuration of hard spheres is a jammed configuration that is the closest one to the initial configuration, hence the term closest jammed configuration.

The paper is structured as follows. Before we present any experimental results, we use Section II to start with definitions that are relevant for the subsequent discussion. These include closest jammed configuration, basin of attraction of a closest jammed configuration, bounding region, bounding surface, and others. We describe the methods that we use to generate packings, to conduct equilibration, and to search for the closest jammed configurations in Section III. Section IV contains the results of packing generation, subsequent equilibration, and searching for the closest jammed configurations. We discuss these results in the same section. Section V presents a summary and conclusions.

II. Definitions

In this section, we briefly provide definitions needed for the rest of the paper. A more elaborate discussion and precise mathematical definitions of most of them can be found in our previous paper.

We rely on the phase space packing description introduced by Salsburg and Wood and therefore use the terms “limiting polytope”, “hypsersurface”, and “hypercylinder” from their paper. Particle velocities are not included in the phase space. Under jamming, we understand collective jamming in packings of frictionless particles. A packing is called jammed if at least a subset of its particles is jammed; other particles are referred to as rattlers. In this paper, we do not exclude rattle particles from packings, when calculating packing densities.

We utilize the concept of inherent structures, initially introduced by Stillinger for packings of particles with soft potential. Inherent structures for such systems are local potential energy minima in the phase space. The potential energy in hard-sphere packings is replaced by the maximum density that a packing can have at a given phase space point, taken with the minus sign. The maximum density is calculated by fixing particle coordinates and inflating particle radii until the first contact between particles occurs. Inherent structures for hard-sphere packings correspond to jammed configurations. To emphasize that we are investigating hard particles, not particles with soft potential, we use below in this paper the term “closest jammed configuration” instead of “inherent structure”.

Each potential energy minimum can be associated with a corresponding basin of attraction, i.e., the energy minimum of a given basin of attraction is the termination point of energy minimization—the steepest descent procedure—for any configuration in this basin of attraction. The bounding region of a given jammed configuration at a given density is by definition the intersection of this configuration’s basin of attraction with the available phase space (when contact hypercylinders for the given density are excluded from the phase space). We define bounding surfaces as surfaces of bounding regions. A bounding region is closed if the bounding surface is fully formed by hypercylinder surfaces.

We define the GCP limit \( \varphi_{\text{GCP}} \) for sufficiently polydisperse packings as the highest possible jamming density of these packings. For monodisperse and slightly polydisperse packings, it is the highest jamming density that can be achieved if crystallization is artificially suppressed. If crystallization is allowed, the GCP limit is revealed by an entropy minimum. We define the RCP limit \( \varphi_{\text{RCP}} \) as the upper bound of the J-segment. Similarly, the lowest typical (LT) jamming density \( \varphi_{\text{LT}} \) is the lower bound of the J-segment.

We distinguish between typical basins of attraction and untypical ones. Basins of attraction with jamming densities in the range \( [\varphi_{\text{LT}}, \varphi_{\text{RCP}}] \) are typical by definition; the others are untypical. With increasing number of particles in the packings, \( \varphi_{\text{RCP}} \) is almost unchanged and \( \varphi_{\text{LT}} \) increases. Thus, we may estimate \( \varphi_{\text{RCP}} \) in the thermodynamic limit by the upper boundary of the J-segment for sufficiently large finite packings. In the present paper, we assume that the J-segment converges in the thermodynamic limit to a single value \( \varphi_{\text{GCP}} \), though this question is still discussed. Let \( \varphi_{\text{max}} \) be the highest possible packing density for a given sphere diameter distribution: it is the crystalline density (~0.74) for monodisperse packings and \( \varphi_{\text{GCP}} \) for sufficiently polydisperse packings. Then, untypical basins of attraction have jamming densities in the range \( [\varphi_{\text{LT}}, \varphi_{\text{LT}}] \cup (\varphi_{\text{RCP}}, \varphi_{\text{max}}) \), where \( \varphi_{\text{LT}} \) is the lowest jamming density, which for monodisperse packings equals at least \( \sqrt{2/\pi}/9 \approx 0.49365 \) (the density of tunneled crystals). Typical and untypical basins of attraction have just been defined for Poisson packings, i.e., when the entire phase space...
is available or, in other words, when initial packing densities are zero. Under typical closest jamming densities for non-zero initial packing densities we understand the closest jamming densities that will be almost always found for packings created at a given density using a given algorithm, if it starts generation at a Poisson configuration. If the jamming density of a bounding region lies in the range $[\phi_{LT}, \phi_{RCF}]$, we refer to this region and all the configurations in this region as liquid. Earlier, we talked about the part of the phase space that is available at a given density. Parts of the available phase space may be completely separated with contact hypercylinders and may have different properties. Thus, it is important to talk about the part of the phase space that is achievable from a given type of configuration. If a bounding region covers almost the entire part of the phase space that is achievable from a given type of configuration, we call this region dominant. Liquid, typical, and dominant basins of attraction coincide for zero initial packing density.

### III. Methods

#### A. Packing generation

Particles in our polydisperse packings have log-normal radii distributions with standard deviations $\sigma$ from 0.05 to 0.3 in steps of 0.05 (the particle mean diameter is normalized to unity). All packings are generated in a fully periodic cubic box and contain $10^4$ particles (cf. Fig. 1). Packings are created in a wide range of compression rates using the force-biased (FB) protocol.\textsuperscript{47–48} This protocol is a modification of the Jodrey–Tory algorithm.\textsuperscript{49–50} The FB algorithm starts from a random distribution of particle centers in a simulation box. Each particle is supplied with an inner diameter chosen to be proportional to the desired particle diameter and to make particles in the closest pair touch each other with their inner diameter shells. Alternatively, a single inner diameter ratio can be specified for the entire packing as the ratio of inner diameters to the desired particle diameters. Similarly, a packing is supplied with an outer diameter ratio, initially larger than unity. The initial outer diameter ratio is chosen to ensure that the total volume of the particles equals the box volume. Particles are also supplied with elastic potential of the third order by overlap distance,\textsuperscript{48} which is cut-off at the outer particle shell. It is now possible to compute the forces between each pair of particles, as well as the net forces for each particle. The algorithm is iterative and each iteration proceeds as follows: (i) determine a net force for each particle; (ii) displace all the particles by distances proportional to the particles’ net forces and in the direction of the net forces; (iii) decrease the outer diameter ratio according to a specified contraction rate; and (iv) update the inner diameter ratio so that the inner diameter shells for the pair of closest particles touch each other. Though the inner diameter ratio may decrease through the iterations, its value has an increasing trend. The algorithm terminates when the outer diameter ratio is equal to the inner diameter ratio. The lower the outer diameter ratio contraction rate, the denser is the final configuration. The source code used in this paper is available under the MIT free software license.\textsuperscript{51}

#### B. Packing equilibration

Salsburg and Wood\textsuperscript{36} derived an equation of state for hard spheres, $p = 1 + 1/[(\varphi_{CJ}/\varphi)^{1/d} - 1]$, where $p$ is the estimated reduced pressure, $\varphi$ is the current packing density, $\varphi_{CJ}$ is the (closest) jamming density for the polytope where the given packing configuration resides, and $d$ is the dimensionality of the system.\textsuperscript{9} The derivation of Salsburg and Wood\textsuperscript{36} assumes that the pressure is stationary and packings are in equilibrium. But all the packings produced in computer simulations or experiments are intrinsically out of equilibrium,\textsuperscript{30,32} because the generation process is non-stationary by definition, especially for fast compressions. The pressure measured in the course of packing generation should therefore not be used for the estimation of $\varphi_{CJ}$ or for the tracking of jamming; instead, packings should be preliminarily equilibrated, \textit{i.e.}, exposed to molecular dynamics simulation with zero compression rate until the pressure is stationary. Equilibration moves the packing to bounding regions that dominate the part of the phase space achievable from an initial configuration.

We equilibrate the packings by performing sets of $2 \times 10^7$ collisions with zero compression rate in a loop until the relative difference of reduced pressures in the last two sets is less than $10^{-3}$, so the pressure can be regarded as stationary. More precisely, to measure the pressure during $2 \times 10^7$ collisions, we average pressures for 100 sub-sets of $2 \times 10^5$ collisions, which amounts to 20 collisions in a sub-set per particle. We use our own implementation\textsuperscript{31} of the Lubachevsky–Stillinger (LS) packing generation algorithm\textsuperscript{53,54} to carry out the equilibration.

#### C. Searching for the closest jammed configurations

To search for the closest jammed configurations, we do not follow the definition of these configurations through the steepest descent energy minimization, but modify the LS algorithm instead. We run the LS algorithm with a high compression rate of 10, until the non-equilibrium reduced pressure reaches a conventional high value of $10^{12}$, then decrease the compression rate by a factor of two and run the LS algorithm again, until the pressure is high enough again ($10^{12}$). We repeat this procedure until the compression rate is $<10^{-4}$. The Boltzmann constant and masses of all particles are set to unity; the temperature is set to 0.2. Fast compressions at the beginning of the search make the initial bounding region collapse as much as possible and at the same time retain the configuration point in this bounding region. Slow compressions at the end of the search allow arriving at a truly jammed configuration. The details and validation of this algorithm are provided in our previous paper.\textsuperscript{14} There we ensured that the distribution of closest jamming densities for packings before equilibration is independent of a particular set of algorithm parameters in a wide range of the latter. We did not repeat this validation for equilibrated packings, as packings before equilibration in the previous paper covered the entire range of polydispersities as well as the entire range of initial and final packing densities considered in the present paper. Searching for the closest jammed configurations after equilibration produces dominant jamming densities.
IV. Results and discussion

In this section, we present the results of the packing equilibration and of searching for the closest jammed configurations of the equilibrated packings. We give an overview of the data in Subsection A (Data overview), analyze the data in Subsection B (Data analysis), test our conclusions for independence from the packing generation protocol in Subsection C (Protocol independence), and finally provide a schematic diagram with the phase space structure in Subsection D (Schematic phase space structure). At the end of the section, in Subsection E (Applicability of liquid equations of state), we check if it is possible to recover the properties of the phase space through comparison of the reduced pressure to predictions from liquid equations of state. To ease the reading of this section, we provide with Table 1 an overview of the symbols used below. Some of them have already been introduced, others will be defined later.

A. Data overview

The dependence of initial packing densities \( \varphi \) on the inverse compression rate \( \gamma^{-1} \) for packings produced with the FB algorithm is shown in Fig. 2a. Even before equilibration, we searched for the closest jamming densities \( \varphi_{\text{CJ}} \) for these packings, as described in Section III C; Fig. 2b depicts for these packings the closest jamming densities \( \varphi_{\text{CJ}} \) vs. initial packing densities \( \varphi \). Then, we equilibrated the initial packings in Fig. 2a and calculated closest jamming density estimates \( \varphi_{\text{JE}} \) from the equation of state by Salsburg and Wood, as described in Section III B. Closest jamming density estimates after equilibration \( \varphi_{\text{JE}} \) vs. initial packing densities \( \varphi \) are shown in Fig. 2c. Finally, we searched for the actual closest jammed configurations of the equilibrated packings (dominant jammed configurations). Dominant jamming densities \( \varphi_{\text{J}} \) vs. initial packing densities \( \varphi \) are shown in Fig. 2d. The same four plots, built vs. \( \gamma^{-1} \), can be found in the Appendix.

We did not average the data in Fig. 2; each point in these figures corresponds to a single packing. To guide the eye, points have been connected by straight lines. Averaging assumes that fluctuations in the data will disappear in the thermodynamic limit. This question is still unresolved.\(^{15,41}\) If fluctuations disappear for infinite packings, averaging also assumes that the noise stemming from finite-size effects is symmetrical around the true value. This is not the case for the closest jamming densities of Poisson packings: the \( \text{J-segment} \) decreases with the increase of the number of particles, but its upper boundary \( \varphi_{\text{RCP}} \) is almost unchanged, only the lower boundary \( \varphi_{\text{LT}} \) is moving upwards.\(^{15} \) Thus, averaging would not produce meaningful results; taking the upper boundary of the \( \text{J-segment} \) instead will give a better estimate of \( \varphi_{\text{RCP}} \) in the thermodynamic limit. Additionally, averaging would remove the information about the exact boundaries of jamming intervals for finite packings.

The RCP densities \( \varphi_{\text{RCP}} \) are included as horizontal lines on the left side of the plots in Fig. 2. Similarly, the GCP densities \( \varphi_{\text{GCP}} \) are shown as horizontal lines on the right side of these plots. We determined the RCP limits as the upper boundaries of the horizontal parts of the plots in Fig. 2b, because the horizontal parts of the plots correspond to the closest jamming densities of Poisson packings.\(^{14}\) We determined the GCP limits for \( \sigma \geq 0.05 \) by asymptotically expanding the plots \( \varphi(1/\gamma) \) (Fig. 2a) and \( \varphi_{\text{CJ}}(1/\gamma) \) (Fig. 7b in the Appendix) into infinite generation time or zero compression rate. \( \varphi_{\text{GCP}} \) for monodisperse packings was taken at the value of a structural transition \( \varphi \approx 0.65 \).\(^{14,22-30} \) More details on the determination of \( \varphi_{\text{RCP}} \) and \( \varphi_{\text{GCP}} \) can be found in our previous paper.\(^{14}\)

### Table 1. Important symbols used in the text

| Symbol | Brief description | Key figures and tables | Values for \( \sigma = 0 \) |
|--------|------------------|------------------------|--------------------------|
| \( \sigma \) | Standard deviation of the log-normal particle radii distributions | X-axis in Fig. 2a and 7 | \(~0.74~) |
| \( \gamma \) | Compression rate for initial packing generation | Y-axis in Fig. 2a, X-axis in Fig. 2b-d | \(~0.64~) |
| \( \varphi \) | Initial packing density after force-biased generation | Y-axis in Fig. 2b | \(~0.49~) |
| \( \varphi_{\text{CJ}} \) | Closest jamming density of a packing | Y-axis in Fig. 2d | \(~0.545~) |
| \( \varphi_{\text{J}} \) | Dominant jamming density of a packing | Left sides of Fig. 2b-d | \(~0.61~) |
| \( \varphi_{\text{LT}} \) | Lowest typical jamming density | Right sides of Fig. 2b-d; Table 2 | \(~0.494~) |
| \( \varphi_{\text{RCP}} \) | Glass close packing limit | \(~0.545~) |
| \( \varphi_{\text{HCP}} \) | Crystalline packing density for monodisperse packings (FCC or HCP crystals) | \(~0.494~) |
| \( \varphi_{\text{max}} \) | Highest packing density: \( \varphi_{\text{HCP}} \) for monodisperse packings, \( \varphi_{\text{GCP}} \) for sufficiently polydisperse packings | \(~0.585~) |
| \( \varphi_{\text{L}} \) | Lowest possible jamming density, at least \( 2/3 \times \varphi_{\text{HCP}} \) for monodisperse packings (density of tunnelled crystals\(^{43}\)) | X-axis in Fig. 2a and d | \(~0.52~) |
| \( \varphi_{\text{m}} \) | Melting transition density (onset of crystallization) | \(~0.585~) |
| \( \varphi_{\text{off}} \) | Offset of crystallization | Fig. 2c and d | \(~0.52~) |
| \( \varphi_{\text{f}} \) | Freezing transition density | Fig. 2c and d | \(~0.52~) |
| \( \varphi_{\text{g}} \) | Ideal glass transition density. \( \varphi_{\text{g}}(\varphi_{\text{g}}) = \varphi_{\text{GCP}} \) | Fig. 2c and d | \(~0.52~) |
| \( \varphi_{\text{MCT}} \) | Density at which available phase space becomes relatively disjoint. \( \varphi_{\text{MCT}} < \varphi_{\text{MCT}} = \varphi_{\text{RCP}} \) | Fig. 2c and d | \(~0.52~) |
B. Data analysis

The closest jamming densities of initial packings (without preliminary equilibration), Fig. 2b, belong to the interval $[4\times 10^{-4}, 4\times 10^{-3}]$ for $\sigma \approx 0.61$ in the case of monodisperse packings and for $\sigma \approx 0.63$ for packings with $s = 0.3$. We may say that for these initial densities liquid bounding regions are typical for the FB algorithm. When we talk about available phase space below in this section, we will always mean available and achievable from the configurations typical for the FB algorithm (which are liquid for $\varphi < [0.61, 0.63]$, the exact value depending on $s$). In the remainder of the paper, we focus on Fig. 2d.

**Spontaneous crystallization, estimation of $\varphi_m$ and $\varphi_{off}$.** For monodisperse packings, bounding regions of configurations with crystalline inclusions dominate the available phase space in the density range $\varphi \approx 0.54$–0.61 ($\sigma = 0$, red line in Fig. 2d). It is manifested by a sudden departure of the $\varphi_J(\varphi)$ plot up to almost crystalline densities. This interval conforms to other studies of spontaneous crystallization and crystal nucleation.\(^5\)–\(^8\) These studies show that the crystal nucleation rate in monodisperse hard sphere packings is negligible for $\varphi < 0.54$, then grows rapidly at $\varphi \approx 0.54$ and reaches a plateau, and then rapidly decreases at $\varphi \approx 0.61.\(^3\) The density $\approx 0.54$ is interpreted as the melting transition density $\varphi_m$ (dedicated studies produce the value $\varphi_m \approx 0.545$ (ref. 1, 4, and 9) for monodisperse packings). The density $\varphi \approx 0.61$ was earlier interpreted as the (ideal) glass transition density $\varphi_g$ for monodisperse hard spheres,\(^5\) which is usually detected by a rapid decrease of compressibility or self-diffusivity. Recent accurate studies show that the ideal glass transition occurs at $\varphi_g \approx 0.585$, while at the same time crystallization is prevented after $\varphi \approx 0.61.\(^4\)–\(^6\) Thus, we refer to $\varphi \approx 0.61$ as the density of the offset of crystallization $\varphi_{off}$. The interval of densities where bounding regions of configurations with crystalline inclusions dominate the phase space of packings with $s = 0.05$ is $\varphi \approx 0.56$–0.58, which can be seen in a sudden departure of the dominant jamming densities in the plot $\varphi_J(\varphi)$ (cyan line in Fig. 2d). This range is also consistent with other results ($\varphi \approx 0.56$–0.59).\(^7\)–\(^9\) Our data report that crystallization becomes impossible for a certain $s$ in the range (0.05, 0.1). Nucleation studies determine that crystallization in packings with Gaussian diameter distribution becomes impossible for $\sigma \approx 0.07,\(^7\)–\(^9\) which conforms to our results (though we investigate log-normal sphere diameter distributions, for such a small polydispersity the two distributions almost coincide).
Change in the phase space structure at $\varphi_{\text{MCT}} \approx 0.52$. For $\varphi \leq 0.52$, the available phase space is strongly connected and dominated by liquid bounding regions for all the packing types. Thus, the closest jamming densities after equilibration are always obtained in the range of liquid jamming densities, $\varphi_l \in [\varphi_{\text{GCP}}^l, \varphi_{\text{MCT}}]$. Starting from a certain density $\varphi \approx 0.52$, the interval of dominant jamming densities moves upward, and starting from a slightly higher characteristic density the dominant jamming density $\varphi_1$ is always higher than $\varphi_{\text{GCP}}$. It means that none of the liquid bounding regions participate in dominating the available phase space any longer. Packings with slight polydispersity $\sigma = 0.05$ clearly demonstrate that the onset of crystallization does not coincide in general with this characteristic density associated with the changes in the structure of the phase space. Changes in the phase space at $\varphi \approx 0.52$, independent of the particle size distribution, are predicted under certain assumptions by the mode-coupling theory; thus, we denote this transition density with $\varphi_{\text{MCT}}$. We assume that bounding regions become relatively disjoint at this density (i.e., the fraction of the “wormholes” in bounding surfaces becomes low).

Glass close packing limit, estimation of $\varphi_{\text{GCP}}$. For packings with $\varphi \geq 0.1$ and $\varphi \geq \varphi_{\text{MCT}}$, the jamming density of the dominant bounding regions increases and reaches maxima at certain densities depending on $\sigma$. We associate these maxima in Fig. 2d with the GCP limits $\varphi_{\text{GCP}}$ (and corresponding packing configurations with the ideal glass), as the GCP limits for packings where crystallization is impossible are by definition the highest jamming densities of these packings. For packings with $\sigma = 0$ and $\sigma = 0.05$ crystallization effects are superimposed on the $\varphi_l(\varphi)$ plots in Fig. 2d. For monodisperse packings crystallization is no longer possible at $\varphi_{\text{MCT}} \approx 0.61$. To the right of this density the dominant jamming density $\varphi_1$ is $\sim 0.65$, which also conforms to the density of the onset of crystalline inclusions in jammed packings and the density where the entropy is minimal $\varphi_{\text{GCP}} \approx 0.65^{14,22-36}$. For polydisperse packings with $\sigma = 0.05$, crystallization is superimposed on the $\varphi_l(\varphi)$ plot in the range $\varphi \approx 0.56-0.58$, but it does not cover the local maximum $\varphi_1 \approx 0.652$ at $\varphi \approx 0.586$; so we attribute $\varphi_1 \approx 0.652$ to the GCP limit of particles with $\sigma = 0.05$. The estimates of the GCP limits compare very well with the GCP limit estimates from our previous paper, where we extrapolated the closest jamming densities of computer-generated packings with $\sigma \approx 0.05$ to infinite generation time. The values from our previous paper are displayed as the lines of corresponding color to the right of Fig. 2d. The differences between the two estimates are $<10^{-2}$ for all the packing types.

Ideal glass transition, estimation of $\varphi_g$. The initial packing density $\varphi$ for which the dominating jamming density reaches its maximum is called the density of the ideal glass transition $\varphi_g$. It is usually measured through the jump in compressibility and divergent alpha-relaxation time. We estimated $\varphi_g$ from Fig. 2d in the following way: (i) first, we took the data points $\varphi_l(\varphi)$ from Fig. 2d in the vicinity of the expected ideal glass transitions (i.e., in the interval $\pm 0.02$ around the global maxima of the $\varphi_l(\varphi)$ plots); (ii) then, we selected local maxima from these data points (because local maxima represent upper boundaries of jamming intervals at each $\varphi_l$; and (iii) fitted these local maxima with third-order polynomials and found the positions of maxima for these polynomials. We consider these positions of maxima as the estimates for $\varphi_g$, which are depicted in Fig. 3. We fitted $\varphi_g$ for $\sigma = 0.05$ with the third-order polynomial and display it in Fig. 3 as well. The extrapolation of this polynomial to monodisperse packings gives a value $\varphi_g = 0.585$. We assume that if crystallization is artificially suppressed in monodisperse packings (e.g., by pinning a certain fraction of particles), the $\varphi_l(\varphi)$ plot will look similar to those for polydisperse packings with $\sigma \approx 0.1$, reaching its maximum value at $\varphi = \varphi_g = 0.585$ with $\varphi_l(\varphi_g) = \varphi_{\text{GCP}}$. It would explain why crystalline inclusions appear in generated packings (prior to equilibration) only at the initial density $\varphi = \varphi_{\text{GCP}} = 0.65^{14,22-36}$ if $\varphi_{\text{GCP}} = 0.65$ is the highest (jamming) density for monodisperse packings with suppressed crystallization, the only way to produce denser packings—for protocols that try to avoid crystallization as long as possible—is to introduce crystalline inclusions in the packing structure. We provide the values for $\varphi_g$ and $\varphi_{\text{GCP}}$ obtained so far in Table 2. Our data comply well with predictions from the mode-coupling theory simulations and experimental observations of divergent relaxation times: $\varphi_g = 0.582$ is reported for packings of $10^3$ particles with uniform distribution having $\sigma = 0.082$; $\varphi_g = 0.59$ is reported for a binary 50 : 50 mixture of $10^3$ particles with diameter ratio 1.4; $\varphi_g = 0.585$ and 0.586 are reported for packings of $2 \times 10^3$ particles with Gaussian distributions of $\sigma = 0.07$ and 0.085, respectively. We display these values in Fig. 3 as well. For binary mixtures, $\sigma$ is computed as the standard deviation of corresponding discrete probabilities of encountering a specific sphere radius.

![Fig. 3](image-url)
We can now call the basins of attraction, the bounding regions, and all the monodisperse and polydisperse packings. This definition conforms to the fact that for \( \sigma \approx 0.1 \) and \( \varphi > \varphi_{\text{MCT}} \), glassy bounding regions dominate the available phase space instead of liquid bounding regions. For monodisperse packings it also conforms to the fact that crystalline inclusions appear in jammed packings only for \( \varphi > \varphi_{\text{GCP}} \). In our previous paper, we referred to the states in closed bounding regions as glassy (following Parisi and Zamponi and Brambilla et al.) because for such states there is only one jammed configuration in the achievable phase space and the configurational entropy is zero. This definition is unsuitable in the light of the current results, as the states in closed liquid bounding regions (that dominate the phase space for \( \varphi \leq \varphi_{\text{MCT}} \)) would be called glassy as well.

C. Protocol independence

To check if our results are protocol-dependent, we did the following: we (i) took the densest jammed packings obtained in Fig. 2 (the FCC crystal for monodisperse packings, the densest partially crystallized packing for \( \sigma = 0.05 \), and the ideal glass packings for \( \sigma \approx 0.1 \)); (ii) proportionally reduced the radii of the particles in these packings to produce unjammed packings in the entire range of densities starting from \( \varphi = 0.4 \); (iii) repeated the procedure for Fig. 2d, i.e., we equilibrated these packings and searched for the closest jammed configurations \( \varphi_{f} \) for the equilibrated packings. Fig. 4 depicts the plot \( \varphi_{f}(\varphi) \) for these diluted densest packings. The horizontal lines to the left and to the right of the figure represent the RCP and GCP limits from Fig. 2d.

Monodisperse packings exhibit a well-known freezing transition at \( \varphi_{\text{f}} = 0.5 \) (the value obtained in dedicated studies is \( \approx 0.494 \)). Polydisperse packings with \( \sigma = 0.05 \) exhibit a freezing transition at \( \varphi_{\text{f}} = 0.52 \). For \( \sigma \approx 0.05 \) and \( \varphi > \varphi_{\text{f}} \), the achievable phase space is dominated by the bounding regions of the densest configurations. For \( \sigma \approx 0.05 \) and \( \varphi \leq \varphi_{\text{f}} \), the achievable phase space is dominated by liquid basins of attraction. The plots in Fig. 2d for \( \sigma \approx 0.05 \) differ from Fig. 4 for \( \varphi \in [\varphi_{\text{f}}, \varphi_{\text{m}}] \cup [\varphi_{\text{off}}, \varphi_{\text{max}}] \). For these \( \varphi \), the bounding regions for the densest configurations dominate the achievable phase space in Fig. 4 but are not achievable from the initial configurations in Fig. 2d.

For polydisperse packings with \( \sigma \approx 0.1 \) and \( \varphi \leq \varphi_{\text{g}} \), the plots in Fig. 4 are qualitatively and quantitatively similar to the ones in Fig. 2d (see the RCP and GCP lines from Fig. 2d in Fig. 4). It means that the conclusions reached in Subsection B (Data analysis) are protocol-independent for these packings; the dominant jamming density does not depend on the initial packing configuration, which determines the achievable portion of the phase space. Thus, we may draw our conclusions in Subsection C for \( \sigma \approx 0.1 \) and \( \varphi \leq \varphi_{\text{g}} \) for the entire available phase space. For \( \varphi > \varphi_{\text{g}} \), the plots differ: the dominant jamming density depends on the initial packing configuration. For diluted densest packings with \( \sigma \approx 0.1 \) and \( \varphi > \varphi_{\text{g}} \), the achievable phase space is dominated by the bounding regions of the ideal glass. It is sometimes argued that the available phase space is fully connected for \( \varphi \leq \varphi_{\text{g}} \) and splits into disconnected portions at \( \varphi_{\text{g}}^{-} \).

These results comply with the general agreement that the available phase space is ergodic at \( \varphi \approx \varphi_{\text{g}}^{-} \). For packings that allow crystallization and for which \( \varphi_{\text{off}} > \varphi_{\text{g}}^{-} \), one may assume that ergodicity breaks at \( \varphi_{\text{off}} \), when the \( \varphi_{f}(\varphi) \) plot becomes protocol-dependent. It was a general assumption in the colloidal literature as well, but recent careful studies of diffusion dynamics show that ergodicity for such packings breaks already at \( \varphi_{\text{g}}^{-} \).

D. Schematic phase space structure

Finally, we present a schematic image with the phase space structure, where we incorporate all the results obtained so far (Fig. 5). We assume that packings that allow crystallization (\( \sigma \approx 0.07 \)) are initially generated in liquid bounding regions for \( \varphi \in [\varphi_{\text{f}}, \varphi_{\text{off}}] \) (as it occurs in Fig. 2d); thus, melting transition and crystallization offset are exhibited for such packings, but not a freezing transition (as it occurs in Fig. 4). Symbols for different

Table 2: Ideal glass transition densities \( \varphi_{\text{g}} \) and corresponding ideal glass densities \( \varphi_{\text{GCP}} \) for packings of hard frictionless spheres with log-normal sphere diameter distributions having different diameter standard deviations \( \sigma \) (as indicated). \( \varphi_{\text{g}} \) vs. \( \sigma \) is plotted in Fig. 3.

| \( \sigma \) | \( \varphi_{\text{g}} \) | \( \varphi_{\text{GCP}} \) |
|---|---|---|
| 0 | 0.585 | 0.65 |
| 0.05 | 0.586 | 0.6518 |
| 0.1 | 0.59 | 0.6549 |
| 0.15 | 0.595 | 0.6596 |
| 0.2 | 0.602 | 0.6645 |
| 0.25 | 0.61 | 0.6711 |
| 0.3 | 0.622 | 0.6779 |
characteristic densities are displayed to the left and below the image. We provide the corresponding values for monodisperse packings to the right and above the image. All the characteristic densities have already been introduced; we only mention that under $\phi_R$ for $\sigma < 0.05$ we understand the ideal glass transition density, $\phi_{GCP}$ is the density of transition from liquid to glass, $\phi_{m}$ is the melting transition density (onset of crystallization), $\phi_g$ is the ideal glass transition density for packings where crystallization is impossible, and $\phi_{off}$ is the density of the offset of crystallization.

The procedure that we used to produce Fig. 2d can be extended to particles with soft potential. In that case, it is necessary to use the particle number density $\rho$ instead of volume density $\phi$. For each number density $\rho$ and temperature $T$, one shall sample the phase space according to the usual equilibrium distribution of states in the canonical ensemble. For each sampled configuration, one shall perform the steepest descent in the potential energy landscape (infinitely fast quenching) and track the value of the obtained potential energy minimum $U$. Presumably, these values will be distributed in one or several narrow intervals around some dominant minima $U(\rho, T)$. There may be more than one of them at a given $\rho$ and $T$ if the phase space is not ergodic. Particle interactions are usually pair-wise; typical model potentials for interactions between soft particles are Gaussian core potential\textsuperscript{58,59} and inverse-power potential.\textsuperscript{60,61} An example for the rapid quenching of several equilibrated monodisperse two-dimensional packings with Gaussian core potential can be found in Stillinger and Weber.\textsuperscript{59} The potential energy of interaction between two particles in inverse-power law systems is $\epsilon(R/r)^{\nu}$, where $r$ is the distance between the two particles and $\epsilon$, $D$, and $n$ are parameters of the potential. Inverse-power law systems have a convenient scaling property. Specifically, it can be shown that all dimensionless excess thermodynamic properties of these systems depend only on a dimensionless reduced number density $\rho_\ast \equiv D^2(\epsilon/\kappa T)^{2/n}$, where $d$ is the packing dimensionality and $k$ is the Boltzmann constant.\textsuperscript{62} For polydisperse packings, $D$ depends on a given particle pair, but it can be split into a dimensionless pair-dependent part and a dimensional pair-agnostic part, of which the latter shall be used for calculating $\rho_\ast$. Thus, the plot $U(\rho, T)$ will be transformed into $U(\rho_\ast)$, where $U_\ast$ is a dominant dimensionless potential energy minimum. We assume that this plot will resemble Fig. 5, only with inverted Y-axis, as far as for hard spheres we defined $U \equiv -\phi_g$. Previous papers on particles with soft potentials typically focused on the distinction between liquid and crystalline phases.\textsuperscript{58,60,61} We believe that finding dominant potential energy minima and building $U(\rho, T)$ plots may shed light on other important properties of such systems, including the ideal glass transition and the distinction between liquid and glass phases.

### E. Applicability of liquid equations of state

At the end of this paper, we investigate for which density ranges liquid equations of state are applicable to the hard-sphere packings under study. We compared reduced pressures $p(\sigma)$ in equilibrated packings (i.e., before densification) from Fig. 4 (diluted densest packings) with values analytically predicted by equations of state for hard spheres. There are many liquid equations of state for polydisperse packings.\textsuperscript{64} We confirm that eqn (9)-(13) from Ogarko and Luding\textsuperscript{64} produce very similar results, as well as eqn (6) from Mansoori et al.\textsuperscript{65} We use the simplest of these equations of state, the one of Boublik–Carnahan–Starling–Mansoori (eqn (4) in Boublik\textsuperscript{63} or eqn (9) in Ogarko and Luding\textsuperscript{64}): $p_{CS}(\sigma) = \frac{1}{1 - \sigma} + O_1 \frac{3\sigma}{1 - \sigma} + O_2 \frac{\phi^3(3\sigma - 1)}{(1 - \sigma)^3}$, where $O_1 = \frac{\langle r \rangle^3}{\langle r^3 \rangle}$ and $O_2 = \frac{\langle r \rangle^2}{\langle r^3 \rangle}$. $\langle r \rangle$ is the $i$th raw moment of the distribution of particle radii $r$. We calculated relative differences $\delta$ between experimental and theoretically predicted reduced pressures, $\delta = \frac{|p(\sigma) - p_{CS}(\sigma)|}{p_{CS}(\sigma)}$, and present in Fig. 6 the $\delta(\sigma)$ plots for different $\sigma$. 

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**Fig. 5** Schematic jamming phase diagram for frictionless hard-sphere packings, depicting dominant jamming density $\phi_d$ vs. initial packing density $\phi$. Opaque areas represent jamming densities that do not dominate the phase space. Red dashed line and red opaque area refer to packings which allow crystallization. Red dashed line and red opaque area refer to packings which allow crystallization. Red dashed line and red opaque area refer to packings which allow crystallization. Red dashed line and red opaque area refer to packings which allow crystallization. Red dashed line and red opaque area refer to packings which allow crystallization.
Naturally, the reduced pressure in all the equilibrated packings at low densities is very close to the theoretical prediction; pressures differ by no more than 1% ($\delta \leq 0.01$). Pressure in packings that allow crystallization ($\sigma < 0.1$) starts to deviate from theoretical predictions at $\varphi_c$. Later on, we will discuss only packings with $\sigma \geq 0.1$, because crystallization effects are not superimposed on their $\delta(\varphi)$ plots. Surprisingly, theoretical predictions for $p(\varphi)$ for these packings remain valid with the same high accuracy $\delta < 0.01$ even for $\varphi > \varphi_{\text{MCT}}$, until $\varphi$ reaches certain values depending on $\sigma$ (Fig. 6). More precisely, $\delta$ is 0.01 at $\varphi \approx 0.581, 0.587, 0.591, 0.596,$ and 0.602 for $\sigma = 0.1, 0.15, 0.2, 0.25,$ and 0.3, respectively. Selecting a different threshold than $\delta = 0.01$ to consider deviations in reduced pressures as high leads to slightly different results. For example, $\delta$ is 0.06 at $\varphi \approx 0.588, 0.596, 0.602, 0.612, 0.622$, for $\sigma = 0.1, 0.15, 0.2, 0.25,$ and 0.3, respectively (Fig. 6). These values of $\varphi$ are very close to $\varphi_g$ from Table 2 for the corresponding $\sigma$. We confirm that the reduced pressure for equilibrated packings from Subsection IV C, cf. Fig. 4. Colors for the different standard deviations $\sigma$ of the log-normal particle radii distributions are depicted in the legend.

Fig. 6  Relative difference $\delta$ between experimental and theoretically predicted reduced pressures vs. initial packing density $\varphi$. $\delta = |p(\varphi) - p_{\text{C.S}}(\varphi)|/p_{\text{C.S}}(\varphi)$. $p_{\text{C.S}}(\varphi)$ is the reduced pressure from the Carnahan–Starling equation of state for polydisperse hard spheres. Experimental pressures $p(\varphi)$ were measured for equilibrated packings from Subsection IV C, cf. Fig. 4. Colors for the different standard deviations $\sigma$ of the log-normal particle radii distributions are depicted in the legend.

Fig. 7  (a) Initial packing densities $\varphi$ vs. the inverse compression rate $\gamma^{-1}$. (b) Closest jamming densities before equilibration $\varphi_{\text{CJ}}$ vs. the inverse compression rate $\gamma^{-1}$. (c) Closest jamming density estimates after equilibration $\varphi_{\text{JE}}$ vs. the inverse compression rate $\gamma^{-1}$. (d) Closest jamming densities after equilibration $\varphi_{\text{JE}}$ vs. the inverse compression rate $\gamma^{-1}$. All the packings were generated with the force-biased (FB) algorithm. Colours for the different standard deviations $\sigma$ of the log-normal particle radii distributions are depicted in the legend.
V. Summary and conclusions

We computer-generated monodisperse and polydisperse frictionless hard-sphere packings of $10^4$ particles with log-normal particle diameter distributions in a wide range of densities $\varphi$ (for monodisperse packings $\varphi = 0.46$–0.72). Then we equilibrated these packings and searched for their closest jammed configurations (inherent structures of hard spheres).

We found that the available phase space is dominated at $\varphi \leq 0.52$ by liquid bounding regions with jamming densities $\varphi_1 \in [\varphi_{LT}, \varphi_{GCP}]$ ($\varphi_{LT} \approx 0.635$ and $\varphi_{GCP} \approx 0.64$ for monodisperse packings). At $\varphi = 0.52$, independent of the particle radii distribution, the structure of the available phase space changes: bounding regions become relatively disjoint (i.e., the fraction of the “wormholes” in bounding surfaces becomes low). The value for this density, also independent of the particle radii distribution, is predicted under certain assumptions by the mode-coupling theory. Thus, we refer to this transition density as $\varphi_{MCT} \approx 0.52$.

For $\varphi > \varphi_{MCT}$, the dominant jamming densities $\varphi_1$ increase with $\varphi$ and the available phase space is dominated by basins of attraction that we call glassy. When $\varphi$ reaches the ideal glass transition density $\varphi_g$, $\varphi_1$ reaches the ideal glass density (the glass close packing limit $\varphi_{GCP}$), so that the available phase space is dominated at $\varphi_g$ by the basin of attraction of the ideal glass. $\varphi_g$ and $\varphi_{GCP}$ depend on the particle size distribution. For packings with sphere diameter standard deviation $\sigma = 0.1$, $\varphi_{GCP} \approx 0.655$ and $\varphi_g \approx 0.59$. For monodisperse and slightly polydisperse packings, crystallization is superimposed on these processes: it starts at the melting transition density $\varphi_m$ and ends at the crystallization offset density $\varphi_{off}$. For monodisperse packings, $\varphi_m \approx 0.54$ and $\varphi_{off} \approx 0.61$. If we extrapolate the ideal glass transition densities $\varphi_g$ for polydisperse packings to $\sigma = 0$ (monodisperse packings), we obtain $\varphi_g \approx 0.585$, in agreement with experiments and simulations on divergent alpha-relaxation time and the jump in compressibility. We verified that the results for packings with $\sigma \approx 0.1$ and $\varphi \leq \varphi_g$ are independent of a packing generation protocol.

We also discovered that the reduced pressure in equilibrated packings complies with liquid equations of state for hard polydisperse spheres for $\varphi > \varphi_{MCT}$. Thus, the comparison with liquid equations of state is not sensitive enough to reveal the changes in the structure of the available phase space at $\varphi_{MCT}$, which are detected by dominant jamming densities.

VI. Appendix

A. Densities vs. inverse compression rate

In Fig. 7, we provide the same plots as in Fig. 2, but built vs. the inverse compression rate $\gamma^{-1}$, not vs. initial packing densities $\varphi$ (as done in Fig. 2b–d). Fig. 7b shows the $\varphi_{GCP}(\gamma^{-1})$ plots that we used to estimate the GCP limits by asymptotic expansion to infinite generation time or zero compression rate. The plots in Fig. 7b have a structure predicted by Parisi and Zamponi\(^4\) (Fig. 2a in that paper).

References

1. G. Parisi and F. Zamponi, Rev. Mod. Phys., 2010, 82, 789–845.
2. S. Khirevich, A. Höltzel and U. Tallarek, Comm. Comput. Phys., 2013, 13, 801–822.
3. S. Khirevich, A. Höltzel, A. Daneyko, A. Seidel-Morgenstern and U. Tallarek, J. Chromatogr. A, 2011, 1218, 6489–6497.
4. M. Skoge, A. Donev, F. H. Stillinger and S. Torquato, Phys. Rev. E: Stat., Nonlinear, Soft Matter Phys., 2006, 74, 041127.
5. L. Filion, M. Hermes, R. Ni and M. Dijkstra, J. Chem. Phys., 2010, 133, 4115.
6. E. Sanz, C. Valeriani, E. Zaccarelli, W. C. K. Poon, P. N. Pusey and M. E. Cates, Phys. Rev. Lett., 2011, 106, 215701.
7. E. Zaccarelli, C. Valeriani, E. Sanz, W. C. K. Poon, M. E. Cates and P. N. Pusey, Phys. Rev. Lett., 2009, 103, 135704.
8. C. Valeriani, E. Sanz, E. Zaccarelli, W. C. K. Poon, M. E. Cates and P. N. Pusey, J. Phys.: Condens. Matter, 2011, 23, 194117.
9. W. G. Hoover and F. H. Ree, J. Chem. Phys., 1968, 49, 3609–3617.
10. L. Berthier and T. A. Witten, Phys. Rev. E: Stat., Nonlinear, Soft Matter Phys., 2009, 80, 021502.
11. G. Brambilla, D. El Masri, M. Pierno, L. Berthier, L. Cipelletti, G. Petekidis and A. Schofield, Phys. Rev. Lett., 2009, 102, 085703.
12. G. Pérez-Ángel, L. E. Sánchez-Diaz, P. E. Ramírez-González, R. Juárez-Maldonado, A. Vizcarra-Rendón and M. Medina-Noyola, Phys. Rev. E: Stat., Nonlinear, Soft Matter Phys., 2011, 83, 060501.
13. T. Voigtmann, A. M. Puertas and M. Fuchs, Phys. Rev. E: Stat., Nonlinear, Soft Matter Phys., 2004, 70, 061506.
14. V. Baranau and U. Tallarek, Soft Matter, 2014, 10, 3826–3841.
15. C. S. O’Hern, L. E. Silbert, A. J. Liu and S. R. Nagel, Phys. Rev. E: Stat., Nonlinear, Soft Matter Phys., 2003, 68, 011306.
16. C. Song, P. Wang and H. A. Makse, Nature, 2008, 453, 629–632.
17. A. Zinchenko, J. Comput. Phys., 1994, 114, 298–307.
18. G. D. Scott and D. M. Kilgour, J. Phys. D: Appl. Phys., 1969, 2, 863–866.
19. Y. Jin and H. A. Makse, Phys. A, 2010, 389, 5362–5379.
20. J. D. Bernal and J. Mason, Nature, 1960, 188, 910–911.
21. J. G. Berryman, Phys. Rev. A, 1983, 27, 1053–1061.
22. A. V. Anikeenko, N. N. Medvedev and T. Aste, Phys. Rev. E: Stat., Nonlinear, Soft Matter Phys., 2008, 77, 031101.
23. A. V. Anikeenko and N. N. Medvedev, Phys. Rev. Lett., 2007, 98, 235504.
24. S. C. Kapfer, W. Mickel, K. Mecke and G. E. Schröder-Turk, Phys. Rev. E: Stat., Nonlinear, Soft Matter Phys., 2012, 85, 030301.
25. B. A. Klumov, S. A. Khrapak and G. E. Morfill, Phys. Rev. B: Condens. Matter Mater. Phys., 2011, 83, 184105.
26. M. Bargiel and E. M. Tory, Adv. Powder Technol., 2001, 12, 533–557.
27. T. Aste and T. Di Matteo, Eur. Phys. J. B, 2008, 64, 511–517.
28. T. Aste and A. Coniglio, Europhys. Lett., 2004, 67, 165–171.
29. K. Lochmann, A. Anikeenko, A. Elsner, N. Medvedev and D. Stoyan, Eur. Phys. J. B, 2006, 53, 67–76.
30. V. Baranau, D. Hlushkou, S. Khirevich and U. Tallarek, Soft Matter, 2013, 9, 3361–3372.
31 P. Chaudhuri, L. Berthier and S. Sastry, Phys. Rev. Lett., 2010, 104, 165701.
32 L. Berthier and G. Biroli, Rev. Mod. Phys., 2011, 83, 587–645.
33 F. H. Stillinger, Science, 1995, 267, 1935–1939.
34 S. Torquato and Y. Jiao, Phys. Rev. E: Stat., Nonlinear, Soft Matter Phys., 2010, 82, 061302.
35 F. H. Stillinger, E. A. DiMarzio and R. L. Kornegay, J. Chem. Phys., 1964, 40, 1564–1576.
36 Z. W. Salsburg and W. W. Wood, J. Chem. Phys., 1962, 37, 798–804.
37 S. Torquato and F. H. Stillinger, Rev. Mod. Phys., 2010, 82, 2633–2672.
38 S. Torquato, T. M. Truskett and P. G. Debenedetti, Phys. Rev. Lett., 2000, 84, 2064–2067.
39 A. Donev, J. Appl. Phys., 2004, 95, 989–999.
40 A. Donev, S. Torquato, F. H. Stillinger and R. Connelly, J. Comput. Phys., 2004, 197, 139–166.
41 M. Pica Ciamarra, M. Nicodemi and A. Coniglio, Soft Matter, 2010, 6, 2871–2874.
42 R. Ni, M. A. C. Stuart and M. Dijkstra, Nat. Commun., 2013, 4, 2704.
43 S. Torquato and F. H. Stillinger, J. Appl. Phys., 2007, 102, 093511.
44 Y. Jiao, F. H. Stillinger and S. Torquato, J. Appl. Phys., 2011, 109, 013508.
45 M. Pica Ciamarra and A. Coniglio, Phys. Rev. Lett., 2008, 101, 128001.
46 M. Pica Ciamarra, P. Richard, M. Schröter and B. P. Tighe, Soft Matter, 2012, 8, 9731–9737.
47 J. Mościński, M. Bargiel, Z. A. Ryczek and P. W. M. Jacobs, Mol. Simul., 1989, 3, 201–212.
48 A. Bezrukov, M. Bargiel and D. Stoyan, Part. Part. Syst. Charact., 2002, 19, 111–118.
49 W. S. Jodrey and E. M. Tory, Phys. Rev. A, 1985, 32, 2347.
50 M. Bargiel and J. Mościński, Comput. Phys. Commun., 1991, 64, 183–192.
51 https://code.google.com/p/packing-generation, 2013.
52 M. C. Vargas and G. Pérez-Ángel, Phys. Rev. E: Stat., Nonlinear, Soft Matter Phys., 2013, 87, 042313.
53 B. D. Lubachevsky and F. H. Stillinger, J. Stat. Phys., 1990, 60, 561–583.
54 B. D. Lubachevsky, J. Comput. Phys., 1991, 94, 255–283.
55 W. van Megen and S. M. Underwood, Phys. Rev. E: Stat., Nonlinear, Soft Matter Phys., 1994, 49, 4206–4220.
56 P. N. Pusey, E. Zaccarelli, C. Valeriani, E. Sanz, W. C. K. Poon and M. E. Cates, Philos. Trans. R. Soc., A, 2009, 367, 4993–5011.
57 R. L. Jack and L. Berthier, Phys. Rev. E: Stat., Nonlinear, Soft Matter Phys., 2012, 85, 021120.
58 F. H. Stillinger, J. Chem. Phys., 1976, 65, 3968–3974.
59 F. H. Stillinger and T. A. Weber, Phys. Rev. A, 1982, 25, 978–989.
60 S. Prestipino, F. Saija and P. V. Giaquinta, J. Chem. Phys., 2005, 123, 144110.
61 C. N. Likos, B. M. Mladek, D. Gottwald and G. Kahl, J. Chem. Phys., 2007, 126, 224502.
62 J. D. Weeks, Phys. Rev. B: Condens. Matter Mater. Phys., 1981, 24, 1530–1535.
63 T. Boublík, J. Chem. Phys., 1970, 53, 471–472.
64 V. Ogarko and S. Luding, J. Chem. Phys., 2012, 136, 124508.
65 G. A. Mansoori, N. F. Carnahan, K. E. Starling and T. W. Leland, J. Chem. Phys., 1971, 54, 1523–1525.