# Soft Matter

## PAPER

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Cite this: DOI: 10.1039/d5sm00379b

Received 14th April 2025, Accepted 27th May 2025

DOI: 10.1039/d5sm00379b

rsc.li/soft-matter-journal

## 1. Introduction

Disordered hyperuniform many-particle systems<sup>1,2</sup> are an emerging exotic class of amorphous states of matter that arise in a variety of contexts and fields, including the eigenvalues of random matrices,<sup>3</sup> nontrivial zeros of the Riemann zeta function,<sup>4</sup> maximally random jammed sphere packings,<sup>5,6</sup> avian photoreceptor mosaics,<sup>7</sup> antigen receptors in the immune system,<sup>8</sup> fermionic ground states,<sup>9,10</sup> nonequilibrium phase transitions,<sup>11-14</sup> active matter,<sup>15</sup> quasicrystals,<sup>16,17</sup> distribution of prime numbers,<sup>18</sup> the large-scale structure of the universe,<sup>19</sup> soft polymeric materials,<sup>20</sup> quantum spin liquids,<sup>21</sup> and myriads of other examples (see ref. 2 and references therein). These correlated disordered systems are characterized by a "hidden order" due to the unusual combination

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Stealthy interactions are an emerging class of nontrivial, bounded long-ranged oscillatory pair potentials with classical ground states that can be disordered, hyperuniform, and infinitely degenerate. Their hybrid crystal-liquid nature endows them with novel physical properties with advantages over their crystalline counterparts. Here, we show the existence of nonequilibrium hard-sphere glasses within this unusual ground-state manifold as the stealthiness parameter  $\chi$  tends to zero that are remarkably configurationally extremely close to hyperuniform 3D maximally random jammed (MRJ) sphere packings. The latter are prototypical glasses since they are maximally disordered, perfectly rigid, and perfectly nonergodic. Our optimization procedure, which leverages the maximum cardinality of the infinite ground-state set, not only guarantees that our packings are hyperuniform with the same structurefactor scaling exponent as the MRJ state, but they share other salient structural attributes, including a packing fraction of 0.638, a mean contact number per particle of 6, gap exponent of 0.44(1), and pair correlation functions  $g_2(r)$  and structures factors S(k) that are virtually identical to one another for all r and k, respectively. Moreover, we demonstrate that stealthy hyperuniform packings can be created within the disordered regime (0 <  $\chi$  < 1/2) with heretofore unattained maximal packing fractions. As  $\chi$ increases from zero, the particles in this family of disordered packings always form interparticle contacts, albeit with sparser contact networks as  $\chi$  increases from zero, resulting in linear polymer-like chains of contacting particles with increasingly shorter chain lengths. The capacity to generate ultradense stealthy hyperuniform packings for all  $\gamma$  opens up new materials applications in optics and acoustics.

> of being statistically isotropic without any long-range order, like ordinary liquids, and yet anomalously suppress infinitewavelength density fluctuations, like perfect crystals and quasicrystals.

> A disordered hyperuniform many-particle system in ddimensional Euclidean space  $\mathbb{R}^d$  is one in which the structure factor  $S(\mathbf{k})$  vanishes as the wavenumber  $k \equiv |\mathbf{k}|$  tends to zero.<sup>1,2</sup> An important subclass are disordered stealthy hyperuniform (SHU) systems in which S(k) = 0 within a spherical "exclusion" region of radius K centered at the origin, *i.e.*, S(k) = 0 for  $0 \le 1$ k < K. SHU many-particle systems are derived as classical ground states of many-particle systems of certain nontrivial, bounded long-ranged oscillatory pair potentials. Remarkably, these hyperuniform ground states can be disordered and infinitely degenerate in the thermodynamic limit.<sup>22-24</sup> The fact that these singular isotropic amorphous states of matter have the character of crystals (S(k) = 0 from infinite wavelengths to intermediate wavelengths of order  $2\pi/K$ <sup>2,25-27</sup> and liquids (statistical isotropy on small length scales)<sup>2</sup> endow them with novel optical, transport, and mechanical properties with advantages over their crystalline counterparts.28-47

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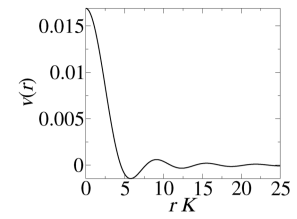
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Existence of nonequilibrium glasses in the degenerate stealthy hyperuniform ground-state manifold



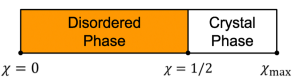
**Fig. 1** The three-dimensional direct-space long-ranged pair potential v(r) versus the dimensionless distance rK associated with the Fourier transform  $\tilde{v}(k)$  that is unity for k < K and zero otherwise, *i.e.*,  $\tilde{v}(k) = v_0\Theta(K - k)$ . We choose  $v_0 = 1$ . This pair potential will be used in our subsequent computations; see eqn (3) with d = 3.

While it is commonplace for quantum-mechanical ground states to be disordered for a variety of typical Hamiltonians,  $^{9,10,48-54}$  it is unusual for classical many-particle systems to remain disordered down to absolute zero temperature with nontrivial interactions, as is the case for disordered SHU ground states. The standard collective-coordinate optimization scheme has been used to create disordered SHU ground states from random initial configurations of *N* particles within a simulation cell with pair potential v(r) under periodic boundary conditions.<sup>22,23,55-57</sup> The total potential energy has the Fourier-space form

$$\Phi(\mathbf{r}^{N}) = \frac{\rho}{2} \sum_{\mathbf{k}\neq\mathbf{0}} \tilde{v}(\mathbf{k}) S(\mathbf{k}), \qquad (1)$$

where  $\tilde{v}(\mathbf{k})$  is the Fourier transform of v(r),  $\rho = N/v_{\rm F}$  is the number density and  $v_F$  is the volume of the fundamental cell. The crucial idea is that if  $\tilde{v}(\mathbf{k})$  is bounded and positive with support in the radial interval  $0 \le |\mathbf{k}| \le K$  and if the particles are rearranged (*via* optimization) so that the structure factor  $S(\mathbf{k})$  is driven to its minimum value of zero for all wavevectors in "exclusion sphere," then the system must be at its ground state or global energy minimum. This class of functions  $\tilde{v}(\mathbf{k})$  lead to bounded longranged oscillatory direct-space pair potentials  $v(\mathbf{r})$ , a threedimensional example of which is shown in Fig. 1. We note that regardless of the choice of  $\tilde{v}(\mathbf{k})$ , the functional form of the potential v(r) is always long-ranged, since  $\tilde{v}(\mathbf{k})$  is a function that has support only in the exclusion region (*i.e.*,  $0 \le |\mathbf{k}| \le K$ ).<sup>24</sup>

SHU ground states have been numerically generated with ultrahigh accuracy.<sup>23,55,56</sup> The disordered ground-state regime for  $d \ge 2$  occurs when  $0 < \chi < 1/2$ ,<sup>24</sup> where  $\chi \equiv M(K)/d(N-1)$  is a dimensionless stealthiness parameter† that specifies the



**Fig. 2** Phase diagram for *d*-dimensional SHU ground states (T = 0) as a function of  $\chi$  for relatively low dimensions with  $d \ge 2$ . The maximal value of  $\chi$ , denoted by  $\chi_{max}$ , depends on the space dimension *d*. Adapted from ref. 24.

number of independently constrained wavevectors, M(K), (within the exclusion sphere of radius K) relative to the total number of degrees of freedom of the system, d(N - 1). ‡ Importantly, the dimensionality of the configuration space per particle,  $d_c$ , decreases linearly with  $\chi$  as  $d_c = d(1 - 2\chi)$  in the thermodynamic limit, explaining the transition from infinitely degenerate disordered phases (when  $\chi < 1/2$ ) to unique crystal structures when  $\chi < 1/2$ ;<sup>24</sup> see Fig. 2. Thus,  $d_c$  is a measure of the cardinality of the infinitely degenerate ground-state manifold set, which, importantly, is maximized in the limit  $\chi \rightarrow 0$ . We will see that this distinguished limit plays a central role in the primary results obtained in this paper.

In the disordered regime, the nature of the energy landscape allows one to find ground states with exquisite precision<sup>23,55,56</sup> with a 100% success rate for relatively large N from random initial conditions. In the limit  $\chi \rightarrow 0$ , *i.e.*, when the cardinality of the infinitely degenerate manifold is maximized, the ground states are ideal-gas-like (Poisson-like), and thus particle pairs can get arbitrarily close to one another. As  $\gamma$  increases from zero, the short-range order and minimum pair separation also increase due to an increase in the number of constrained degrees of freedom in a finite-N system.<sup>23,24,55</sup> Such high- $\chi$ disordered ground states have been generated to create SHU sphere packings by circumscribing the points by identical nonoverlapping spheres.58 However, the success rate to find allowable packing configurations among all ground states obtained with even moderate values of the packing fraction  $\phi$ (<0.25) falls off rapidly with N and vanishes in the thermodynamic limit.59

Is it possible to exploit the huge degeneracy of the energy landscape to obtain much denser stealthy ground-state packings by biasing the search in configuration space so that such atypical portions of the manifold are found? In this article, we show this possible and, by doing so, shed new light on the nature of the infinitely degenerate ground-state manifold under the action of a generalized stealthy long-ranged potential (defined in eqn (2)). We begin by demonstrating that this manifold, counterintuitively, in the zero-stealthy limit ( $\chi \rightarrow 0$ ) when the cardinality is maximized, contains states that are remarkably configurationally extremely close to the hyperuniform nonequilibrium MRJ sphere packings that were recently reported by Maher *et al.*<sup>6</sup> using the linear programming

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<sup>†</sup> In the thermodynamic limit,  $\chi$  is inversely proportional to  $\rho$  according to the exact relation<sup>24</sup>  $\rho\chi = \nu_1(K)/[2d(2\pi)^d]$ , where  $\nu_1(R)$  is the volume of a *d*-dimensional sphere of radius *R*.

<sup>‡</sup> Although there are 2M(K) + 1 wavevectors inside the spherical exclusion region of radius *K* centered at the origin, only M(K) of them are independently constrained because  $S(\mathbf{k})$  is inversion symmetric, *i.e.*,  $S(\mathbf{k}) = S(-\mathbf{k})$ .

packing algorithm of Torquato and Jiao.<sup>60</sup> § The MRJ packing state is a prototypical nonequilibrium glass, since it is the most disordered packing subject to strict jamming,¶ resulting in packings that are perfectly nonergodic (i.e., permanently trapped in configuration space) and possess infinite elastic moduli.<sup>67,68</sup> For this reason, it is remarkable the aforementioned special ground-state packings and MRJ packings share the following salient structural attributes: pair correlation functions  $g_2(r)$  and structures factors S(k) that are virtually identical to one another for all r and k, respectively, a packing fraction of 0.638, a mean contact number per particle of 6 and a "gap" exponent of 0.44. Moreover, our 3D ground-state packings become hyperuniform, but appropriately no longer stealthy, in the thermodynamic (infinite-size) limit with the same structure-factor scaling exponent as the MRI state. Figuratively, this is akin to finding a nonequilibrium "needle" in a "haystack."

We also show that a large family of other stealthy hyperuniform packings can be created as  $\chi$  increases up to 1/2 with heretofore unattained maximal packing fractions. These packings also have interesting structural characteristics; the particles in this family of disordered packings always form interparticle contacts, albeit with sparser contact networks as  $\chi$  increases from zero, resulting in linear polymer-like chains of contacting particles that progressively possess shorter mean chain lengths. (The reader is referred to ref. 59 for a comprehensive study of SHU packings with and without soft-core repulsions within the disordered regime for d = 1, 2, 3). This capacity to generate ultradense SHU packings opens up new applications in optics and acoustics relative to previous studies that used low-density SHU packings.<sup>29,34,36,43,69</sup>

The rest of the paper is organized as follows: in Section 2, we describe the modified collective-coordinate optimization scheme and how it is applied to determine the maximal packing fraction of SHU packings with a given value of  $\chi$ . Here we also describe how to determine whether the corresponding contact networks percolate. In Section 3, we provide results for the ultradense SHU packings with  $\chi$  tending to zero and select positive values of  $\chi$  within the disordered regime, including their maximal packing fractions, pair statistics, and mean contact numbers. Finally, in

Section 4, we provide concluding remarks and outlook for future work.

### 2. Simulation method

### 2.1. Generation of dense disordered SHU packings

To achieve the desired goal of creating ultradense disordered SHU packings, we modify the total potential energy function (1) by including an additional pairwise soft-core repulsive potential u(r) that is bounded, differentiable, and positive with support in the finite range  $0 \le r < \sigma$ :<sup>36,57,69</sup>

$$\Phi(\mathbf{r}^{N}) = \frac{\rho}{2} \sum_{\mathbf{k}\neq\mathbf{0}} \tilde{v}(\mathbf{k}) S(\mathbf{k}) + \sum_{i < j} u(r_{ij}).$$
(2)

We choose  $\tilde{v}(k) = v_0 \Theta(K - k)$ , where  $\Theta(x)$  is the Heaviside step function, yielding the bounded long-ranged pair potential

$$\frac{v(r)}{v_0} = \left(\frac{K}{2\pi r}\right)^{d/2} J_{d/2}(Kr),$$
(3)

where  $J_{\nu}(x)$  is the Bessel function of the first kind of order  $\nu$ . The soft-core contribution to the energy (2), embodied by u(r), competes with standard stealthy contribution, and hence the modified potential biases the optimization to sample portions of the SHU ground-state manifold in which the minimum particle pair separation is substantially greater than those obtained via the stealthy contribution only. Since both sums in (2) are nonnegative, its ground states with  $\Phi(\mathbf{r}^N) = 0$  must satisfy the SHU condition [*i.e.*, S(k) = 0 for  $0 \le k < K$ ] such that all particle pairs are separated by at least a targeted distance  $\sigma$ . Then, such ground states are mapped to SHU packings with a packing fraction at least given by  $\phi = \rho v_1(\sigma/2)$ , where  $v_1(a) = \pi^{d/2} a^d / \Gamma(1 + d/2)$  is the volume of a *d*-dimensional sphere of radius *a*. At a specific value of  $\chi$ , the packing fraction  $\phi$  can be increased to its maximum value,  $\phi_{\max}(\chi)$ , beyond which the ground state ceases to exist, which can be viewed as a satisfiable-unsatisfiable (SAT-UNSAT) transition.<sup>70,71</sup> We numerically determine the values of  $\phi_{\max}(\chi)$ , as described in Section 2.2. The modified potential (2) was previously used to generate the SHU ground-state sphere packings but only at relatively small packing fractions for optical and elastodynamic applications.36,69

A general form of the soft-core repulsion u(r) in the potential energy (2) is

$$\frac{u(r)}{\varepsilon_0} = \left(1 - \frac{r}{\sigma}\right)^{\beta} \Theta(\sigma - r) \quad (\beta > 1).$$
(4)

In this work, we set  $\beta = 2$ , yielding the harmonic contact potential,<sup>61,72,73</sup> because it is computationally efficient to implement.

Importantly, the ground states of potential (2) are independent of the ratio of  $v_0/\varepsilon_0$  if  $v_0$  and  $\varepsilon_0$  are positive and finite. We consider a cubic fundamental cell, and set the energy scales to be  $v_0 = 1$  and  $\varepsilon_0 = 100v_0$ , enabling us to generate valid packings that are ground states in a reasonable amount of computational time for  $N \approx 400-10774$  and  $n_c = 10^2-10^4$ , where  $n_c$  is the number of configurations. Specifically, using an Intel(R)

<sup>§</sup> Such MRJ-like disordered jammed states have been created by various protocols and systems under periodic boundary conditions, including rapid compression of hard spheres<sup>5,6,60,61</sup> to their deep mechanically stable local "energy" ( $-\phi$ ) minima (inherent structures<sup>60,62</sup>), rapid quenching of soft spheres at high *T* to find inherent structures at *T* = 0 *via* conjugate gradient techniques,<sup>61,63</sup> or as a dynamical phase transition *via* a biased random organization protocol.<sup>64</sup> It is well-known that these current packing protocols lead to disordered jammed packings with similar but not exactly the same structural features, such as rattler concentrations, gap exponents, hyperuniformity, and force distributions. For example, the Torquato–Jiao linear programming algorithm<sup>60</sup> produces 50% less rattlers<sup>65</sup> than that of the modified Lubachevsky-Stillinger algorithm.<sup>66</sup> Work on rapid quenching of soft spheres<sup>61,63</sup> typically use only the soft-core potential in (2), *i.e.*, without the stealthy contribution, and hence cannot guarantee hyperuniformity of the packing.

<sup>¶</sup> A packing is strictly jammed if there are no possible collective rearrangements of some finite subset of particles, and no volume-nonincreasing deformation that can be applied to the packing without violating the impenetrability constraints of the particles.<sup>68</sup>

Xeon(R) CPU (E5-2680, 2.40 GHz), it takes approximately 15 core-hours to generate one 3D ground state with  $\chi = 0.45$ ,  $\phi = 0.47$ , and N = 4000. It takes roughly 130 core-hours on the same CPU to generate one 3D ground state with  $\chi = 9.28 \times 10^{-5}$ , N = 10774, and  $\phi = 0.638$ .

The algorithm to find ultradense SHU ground-state packings is performed as follows: For given parameters of system size N,  $\chi(>0)$ , and target packing fraction  $\phi = \phi_{max}(\chi)$ , we begin with a random initial condition in a simple cubic fundamental cell and minimize its potential energy  $\Phi$  given in eqn (3) *via* the low-storage Broyden–Fletcher–Goldfarb–Shanno (L-BFGS) algorithm.<sup>74,75</sup> The minimization stops when (i)  $\Phi < 5 \times 10^{-20}$ , || (ii) the number of evaluations exceeds  $5 \times 10^6$ , or (iii) the mean particle displacements are less than  $10^{-15}\rho^{-1/d}$ . After terminating the minimization, we retain the ground-state point pattern that satisfies condition (i) and discard it otherwise. The values of system size N, the number of configurations  $n_c$ , and stealthiness parameter  $\chi$ are listed in Table 3.

### 2.2. Determination of maximum packing fractions

We first determine the maximal target packing fraction of 3D SHU ground states with the smallest value of  $\chi$ , denoted by  $\chi_{min}$ . For the simple cubic fundamental cell we consider,  $\chi_{min}$  is a function of particle number *N*:

$$\chi_{\min}(N) = \frac{1}{N-1},\tag{5}$$

where we have used the definition of  $\chi$ , the fact that the smallest number of constraints is M(K) = 3 with  $K = 2\pi/L$ , and L is the side length of the fundamental cell. The explicit formula for the total potential with  $\chi = \chi_{\min}(N)$  is provided in eqn (9) in Appendix B. Thus,  $\chi_{\min}$  vanishes as N tends to infinity. At each system size N, we numerically determine  $\phi_{\max}(\chi_{\min})$  as follows: For a target packing fraction  $\phi$ , we carry out the optimization mentioned above (Section 2.1) for either a given number of initial conditions  $n_i$  (=100) or at most 1680 corehours, equivalent to one week for 10 threads of an Intel(R) Xeon(R) CPU (E5-2680, 2.40 GHz). The success ratio is obtained by dividing the number of ground states by the number of used initial conditions. This search is done by varying  $\phi$  from 0.630 to 0.640 in increments of 0.001. The values of  $\phi_{\text{max}}(\chi)$  are determined as the largest  $\phi$  such that the success ratio is at least 7%. Using the determined value of  $\phi_{\max}(\chi)$  (see Table 1), we then generate  $n_{\rm c}$  independent ground states (as tabulated in Table 3).

Using the same procedure, we also numerically determine  $\phi_{\max}(\chi)$  for  $\chi = 0.10, 0.20, 0.30, 0.35, 0.40$ , and 0.45, but apply a slightly looser criterion than for  $\chi_{\min}$  to save overall computation cost in both determination of  $\phi_{\max}(\chi)$  and generation of the ultradense SHU packings. At each  $\chi$  value, we use either a given number of initial conditions  $n_i(=50)$  or, at most, 840 core hours to obtain ground states. We consider the largest value of  $\phi$  achieved at least 10% of success ratio as  $\phi_{\max}(\chi)$ . This search is done by varying  $\phi$  from 0.20 to the packing fraction of the

**Table 1** Maximal packing fractions  $\phi_{max}(0^+)$  and fitted values of the gap exponent  $\gamma$  for our ultradense SHU ground-state packings with  $\chi = 0^+$ , and 3D MRJ packings. Data for the MRJ packings were obtained in ref. 6. The values of  $\chi_{min}$  are given in eqn (5). The values in parentheses stand for the statistical errors

Models	Ν	$\phi_{\max}\left(0^{^{+}} ight)$	γ
SHU packings	400	0.636	0.45(1)
SHU packings	1198	0.637	0.44(1)
SHU packings	3592	0.637	0.44(1)
SHU packings	10774	0.638	0.44(1)
3D MRJ packing	5000	0.638	0.44(1)

densest lattice packing in increments of 0.01. The obtained values of  $\phi_{\max}(\chi)$  are tabulated in Table 2.

#### 2.3. Percolation of contact networks

For the contact networks explained in the caption of Fig. 4, we determine whether they percolate by a simplified version of the algorithm for general networks, explained in ref. 58 and 76. Specifically, we begin by creating a contact network of a configuration in which each vertex (or node) is a particle and an edge is formed between any two particles whose separation is less than  $\sigma + 10^{-3} \rho^{-1/3}$ . For such a network, we randomly choose an initial vertex and recursively search for all vertices connected to it. This search stops under two conditions: (i) when there are no further vertices to explore or (ii) when a newly searched vertex is a periodic copy of the initial vertex. When condition (ii) is met, we terminate the search for that configuration and consider the associated contact network to be percolated. Otherwise, we repeat this procedure up to 1000 different initial vertices for each configuration. Table 2 of the Results section provides the percentage of contact networks that percolate, *i.e.*, topologically connected across the sample.

## 3. Results

Employing our modified collective-coordinate optimization procedure described in Section 2, we demonstrate that our disordered packings, as  $\chi$  tends to zero, become hyperuniform and non-stealthy in the thermodynamic limit with the same structure-factor scaling exponent as the MRJ state reported in

**Table 2** Summary of parameters of 3D disordered sphere-packing ground states across stealthiness parameter  $\chi$ . Three parameters include maximal packing fractions  $\phi_{max}(\chi)$ , mean contact number per particle  $Z(D^+)$ , and the fraction of configurations that have percolated contact networks *p*. For other values of  $\chi$ , we consider 1000 ground states with N = 4000 taken from ref. 59. The values in the parentheses for  $Z(D^+)$  indicate statistical errors

χ	$\phi_{\max}(\chi)$	$Z(D^+)$	p [%]
0+	0.638	6.0(1)	100.0
0.10	0.61	4.388(4)	100.0
0.20	0.58	2.960(4)	100.0
0.30	0.55	1.975(3)	100.0
0.35	0.53	1.514(3)	38.6
0.40	0.50	0.975(23)	0.0
0.45	0.47	0.600(21)	0.0

 $<sup>\</sup>parallel$  This threshold is close to zero energy of the potential, given in eqn (3), within the double precision of the machine

**Table 3** Simulation parameters for 3D ultradense SHU ground-state packings at  $\chi$ . Here, *N* is the system size, and  $n_c$  is the number of independent ground states

Ν	χ	n <sub>c</sub>
400	$\chi_{\rm min} = 2.51 \times 10^{-3}$	10 000
1198	$\gamma_{\rm min} = 8.35 \times 10^{-4}$	5000
3592	$\chi_{\rm min} = 2.78 \times 10^{-4}$	1000
10774	$\chi_{\rm min} = 9.28 \times 10^{-5}$	100
4000	0.10	500
4000	0.20	500
4000	0.30	500
4000	0.35	500
4000	0.40	500
4000	0.45	500

ref. 6. Furthermore, they share other salient structural attributes, including a packing fraction of 0.638, a mean contact number per particle of 6 (isostaticity), a gap exponent  $\gamma = 0.44$ , and pair correlation functions  $g_2(r)$  and structure factors S(k) that are virtually identical to one another for all r and k, respectively. We begin by showing that we can create other stealthy hyperuniform packings whose particles always form interparticle contacts for positive values of  $\chi$  as it increases up to 1/2 with heretofore unattained high packing fractions and determine their maximal packing fractions as well as mean contact numbers per particle. Subsequently, we focus on the pair statistics of the hyperuniform packings in the limit  $\chi \rightarrow 0$  and SHU packings with  $\chi = 0.45$ . We determine the limit  $\chi \rightarrow 0$  by applying  $\chi = \chi_{\min}(N)$  given by eqn (5) and the associated total potential  $\Phi$  given in eqn (9) for a given N to a sequence of increasing system sizes with N = 400, 1198, 3592 and 10774, yielding  $\chi_{\rm min} = 2.51 \times 10^{-3}$ ,  $8.35 \times 10^{-4}$ ,  $2.78 \times 10^{-4}$  and  $9.27 \times 10^{-5}$ , respectively, and then extrapolating to the limit  $\chi = 0$  by increasing *N*. Importantly, the potential (9) guarantees the obtained ground states are (stelathy) hyperuniform for any N, different from compression algorithms. Henceforth, we will refer to this limit simply as  $\chi = 0^+$ .

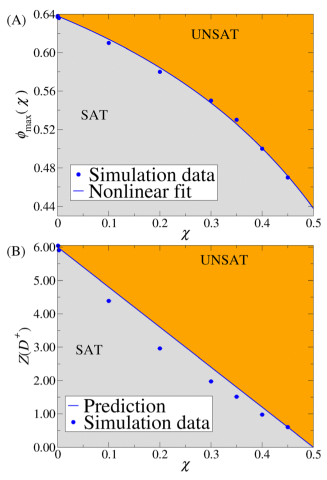
#### 3.1. Maximal packing fractions and mean contact numbers

Application of our modified collective-coordinate optimization procedure (see Section 2 for details) within the disordered regime (0 <  $\chi$  < 1/2) yields achievable packing fractions that are substantially larger than those that can be obtained from the standard collective-coordinate procedure without the soft-core repulsive potential. Importantly, the maximal packing fraction decreases (not increases) as  $\chi$  increases and approaches  $\chi = 1/2$ , as seen in Table 1 and Fig. 2(A). The fact that  $\phi_{\text{max}}$  is highest in the small- $\chi$  limit and then monotonically decreases with  $\chi$  up to  $\chi = 1/2$  is consistent with the decrease in the relative degrees of freedom as  $\chi$  increases. Such  $\chi$ -dependence on  $\phi_{\text{max}}(\chi)$  can be well approximated by the following [1,1] Padé approximant:

$$\phi_{\max}(\chi) = 0.638 \frac{1 - 1.245 \chi}{1 - 0.900 \chi},\tag{6}$$

which we see is in very good agreement with the data plotted in Fig. 3(A).

It is noteworthy that for all  $\chi$  values within the disordered regime, particles in this family of packings always form



**Fig. 3** Parameters of 3D disordered ultradense SHU sphere-packing ground states across stealthiness parameter  $\chi$ . (A) Maximal packing fraction  $\phi_{max}(\chi)$  as a function of  $\chi$ . The filled circles are simulation data (with *N* up to 10774 and  $10^2-10^4$  configurations) and the solid curve is the [1,1] Padé approximant of the data given by eqn (6). (B) Mean contact number per particle  $Z(D^+)$  as a function of  $\chi$ . The solid line is the theoretical prediction (8). The data in both (A) and (B) are tabulated in Table 2 in the Results section. Note that the gray and orange regions in (A) and (B) correspond to the SAT and UNSAT phases<sup>70,71</sup> of the ground state of eqn (2), respectively. The reader is referred to ref. 59 for details.

interparticle contacts. The quantity

$$Z(r) = 4\pi\rho \int_0^\infty r^2 g_2(r) \mathrm{d}r \tag{7}$$

is the cumulative coordination number from which we can extract the mean contact number per particle,  $Z(D^+)$ , and we take  $D^+$ (indicating the limit of *r* to *D* from above) to be  $\sigma_{\text{max}} + 0.001 \rho^{-1/3}$ . As shown in both Fig. 3(B) and Table 2, the mean contact number per particle is the largest as  $\chi$  tends to zero, where  $Z(D^+)$  achieves the isostatic (marginally mechanically stable) value of 6 associated with strictly jammed packings and then monotonically decreases with  $\chi$ , implying sub-isostatic packings ( $Z(D^+) < 6$ ) that cannot be strictly jammed.<sup>68</sup> Such a decrease in  $Z(D^+)$  with  $\chi$  is due to an interplay between two competing constraints in numerical optimizations. Specifically, for an *N*-particle SHU ground-state packing in  $\mathbb{R}^d$ ,  $2\chi \times d(N - 1)$  degrees of freedom are already constrained among d(N - 1) total degrees of freedom by the stealthy hyperuniform condition. Thus, the remaining degrees of freedom determine the maximum number of constraints from effectively contacting particles:

$$Z(D^{+}) = 2d(1 - 2\chi).$$
(8)

This prediction shows good agreement with the simulation data; see Fig. 3(B).

Fig. 4 shows representative contact networks of the ultradense SHU packings at selected values of  $\chi$  between 0 to 1/2, some of which percolate across the entire system. At  $\chi = 0^+$ , shown in Fig. 4(A), the contact network percolates and effectively satisfies the isostatic condition  $Z(D^+) \approx 6.0$ , implying that these packings are effectively jammed. As  $\chi$  increases from zero, the contact networks are always sub-isostatic, as predicted by

eqn (8). For  $\chi$  = 0.20, the contact network percolates, and almost all of the spheres are part of it, but the network becomes sparser (contact number decreases) from the isostatic value of 6 for  $\chi \to 0$  with  $Z(D^+) \approx 3.0$ , as seen in Fig. 4(B) and Table 2. As  $\chi$  increases from 0.20 to 1/2,  $Z(D^+)$  continues to decrease, *i.e.*, the contact networks become even sparser, resulting in linear polymer-like chains of contacting particles that progressively possess shorter mean chain lengths. This trend is evident in Fig. 4(C) for  $\chi = 0.35$  with  $Z(D^+) \approx 1.5$ , and in Fig. 4(D) for  $\chi =$ 0.45 with  $Z(D^+) \approx$  0.6. In addition, approximately 38.6% of the contact networks percolate when  $\chi = 0.35$ , and they then cease to percolate for the higher values of  $\chi$ ; see Table 2 and Section 2.3 for details about how we estimate percolation behaviors. When  $\chi = 0.45$ , the non-percolating contact network consists of dimer polymer-like chains. Here, about half of the particles in the packing are monomers. The reader is referred to ref. 59 for

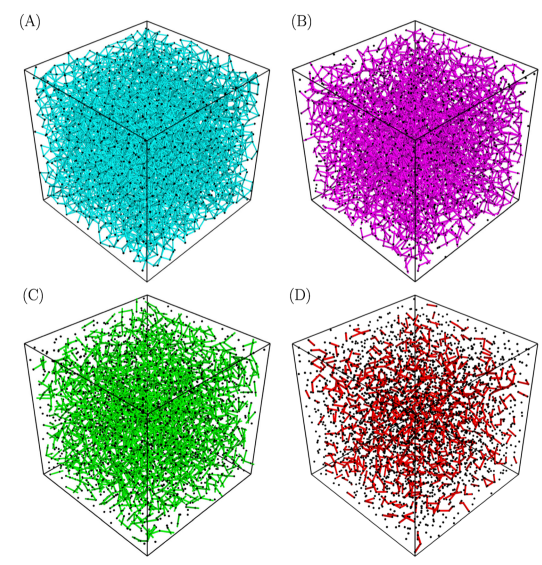


Fig. 4 Contact networks of 3D SHU sphere packings for various  $\chi$  values: (A)  $\chi = \chi_{min}(N)$ , N = 3592,  $\phi = 0.637$  and  $Z(D^+) \approx 6.0$ ; (B)  $\chi = 0.20$ , N = 4000,  $\phi = 0.58$ , and  $Z(D^+) \approx 3.0$ ; (C)  $\chi = 0.35$ , N = 4000,  $\phi = 0.53$ , and  $Z(D^+) \approx 1.5$ ; (D)  $\chi = 0.45$ , N = 4000,  $\phi = 0.47$  and  $Z(D^+) \approx 0.60$ . Here, the sizes of the cubic frames are identical. The centroids of the particles are depicted as black dots in all panels. In each panel, the bonds are drawn between particles with colored lines if pairs of particle centers are closer than  $\sigma + 0.001\rho^{-1/3}$ .

the details behind these results for all  $\chi$  as well as corresponding findings concerning local spatial statistics (nearestneighbor and minimal pair-distance distributions) for not only 3D cases, but 1D and 2D instances.

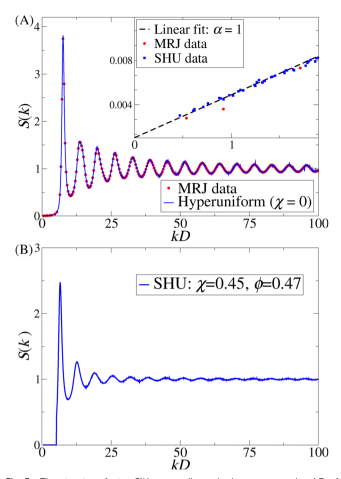
### 3.2. Structure factors

An isostatic value of  $Z(D^+) = 6.0$  and packing fraction  $\phi = 0.638$ are not sufficient to define an MRJ packing. Therefore, we now show that the ground-state packings in the limit  $\chi \to 0$  (*i.e.*, when the cardinality of the infinitely degenerate ground-state manifold) are configurationally extremely close to the MRJ state, as measured by structure factors S(k) that are virtually the same as one another for all k. While MRJ packings are hyperuniform, they are not perfectly stealthy,<sup>6</sup> and hence the limit  $\chi \rightarrow 0$  (corresponding to maximal cardinality of the infinite set of degenerate ground states) is a necessary condition to check if SHU packings have any correspondence to the MRJ state, since the stealthiness vanishes in this limit while being hyperuniform, which importantly implies the thermodynamic (infinite-N) limit. To approach the large-N limit, we study a sequence of increasing systems sizes with N = 400, 1198,3592 and 10774, while taking  $\chi = \chi_{\min}(N)$  given by eqn (5) and then extrapolate to the limit  $N \rightarrow \infty$ . We find that as  $\gamma$  becomes small, the already small stealthy region diminishes in size, and remarkably, the ground states become hyperuniform but not stealthy in the limit  $\chi \to 0$ . We also confirm that the ultradense SHU packings with  $\chi = \chi_{min}$  exhibit the power-law scaling form  $S(k) \sim k$  as  $k \to 0$  by extrapolating S(k) in the range of KD < kD  $\lesssim$  1.3, which is consistent with those for MRJ state reported in ref. 6; see the inset of Fig. 5(A). Fig. 5(A) also compares the structure factor for a wide range of wavenumbers for the SHU packings at the largest value of N (=10774) to the recent corresponding MRJ data,6 revealing that they are virtually identical to one another on the scale of this figure.

By contrast, the structure factors S(k) of the SHU spherepacking ground states with the larger values of  $\chi$  are quite distinct from those in the zero- $\chi$  limit; see Fig. 5(B) for the case  $\chi = 0.45$ . When the ground states have large values of  $\chi$ , S(k)clearly have a wide exclusion region in which S(k) = 0 for small k. Furthermore, the oscillations in S(k) decay more rapidly as wavenumber k increases, reflecting a decrease in the degree of order at short-range length scales due to the formation of dimer polymer-like chains.

# 3.3. Pair correlation functions and cumulative coordination number

To further confirm the striking correspondence to MRJ packings, we compare its pair correlation function  $g_2(r)$  to that of our ground-state packings with  $\chi = 0^+$  in Fig. 6(A) for the largest system size. Again, we see that the pair correlation functions are virtually identical to one another on the scale of this figure, including subtle small-scale structural features. Specifically, they share a Dirac-delta-like peak at r = D, the well-known split second peak seen in disordered jammed packings at  $r/D = \sqrt{3}$ and 2, as well as the power-law singularity for near contacts.<sup>66</sup>

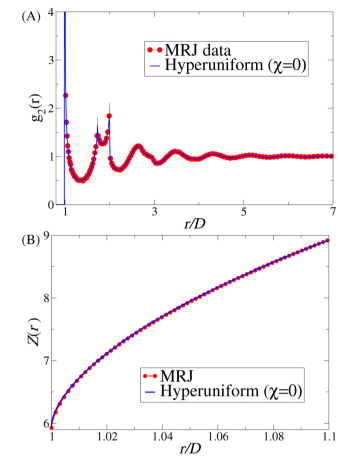


**Fig. 5** The structure factor *S*(*k*) *versus* dimensionless wavenumber kD of 3D SHU sphere-packing ground states. (A) Comparison of the data of 3D hyperuniform MRJ sphere packings from ref. 6 to the corresponding structure factor for our 3D hyperuniform sphere-packing ground states in the limit  $\chi \rightarrow 0$ , as extrapolated from the various system sizes studied, namely, N = 400, 1198, 3592, 10774. Both models have the same packing fraction,  $\phi = 0.638$ . The inset shows *S*(*k*) in the range of kD < 2 for SHU sphere-packing ground states with  $\chi = \chi_{min}(N)$  as obtained for the aforementioned sizes and MRJ states. The black dashed line is a linear fit of the data, *i.e.*, *S*(*k*) ~ *k* in the limit  $k \rightarrow 0$ . (B) 3D SHU sphere-packing ground states with  $\chi = 0.45$ , N = 4000,  $\phi = 0.47$ , and  $n_c = 500$  are shown. In both (A) and (B), *D* is the hard-sphere diameter.

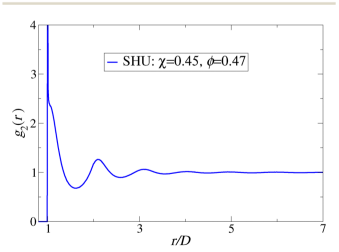
For general dense packings, the exponent  $\gamma$  in the power-law singularity, *i.e.*,  $g_2(r) \sim (r/D - 1)^{-\gamma}$  for 0 < r/D - 1 < 0.2, is expected to increase with increasing short-range order.<sup>66</sup> Specifically,  $\gamma \rightarrow 0$  means a constant  $g_2$  near contact, which is a signature of the ideal gas. In contrast,  $\gamma \rightarrow 1$  means that the first and second peaks are clearly separated with a wide range of gaps, which is typical of crystal packings. Thus,  $\gamma$  for the MRJ-like SHU ground states should lie between 0 and 1.

The excellent agreement of the near-contact divergence in the ground-state packings with  $\chi = 0^+$  and MRJ packings is more apparent in the cumulative coordination number Z(r); see Fig. 6(B). Here, we find that both packings yield the same fit  $Z(x) \sim Z_{\text{fit}} + Cx^{1-\gamma}$ , where  $x \equiv r/D - 1$ . In the near-contact region (*i.e.*, 0.001 < x < 0.1), the fitted gap exponents are  $\gamma = 0.44(1)$  for largest ultradense SHU packings with  $\chi = \chi_{\text{min}}$  and the MRJ

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**Fig. 6** Comparisons of 3D hyperuniform sphere-packing ground states ( $\chi = 0^+$ ) and 3D MRJ sphere packings in direct space. (A) The pair correlation function  $g_2(r)$  versus dimensionless distance r/D of the two models. (B) The cumulative coordination number Z(r) versus dimensionless distance r/D of the two models. In both panels, we consider 3D MRJ packings taken from ref. 6 and D is the hard-sphere diameter. In both (A) and (B), two models have the same packing fraction,  $\phi = 0.638$ .



**Fig. 7** The pair correlation function  $g_2(r)$  versus dimensionless distance r/D of 3D SHU sphere-packing ground states with a high value of  $\chi$ . Here, D is the hard-sphere diameter. We consider the ground states with  $\chi = 0.45$ , N = 4000,  $\phi = 0.47$ , and  $n_c = 500$ .

state. Table 1 summarizes the values of  $\gamma$  at various system sizes *N*. Interestingly, by the same contact criterion stated in the caption of Fig. 4, we estimate the rattler concentration to be about 2%, which is close to the recently reported value for MRJ packings.<sup>6</sup>

For the SHU packings with the largest value of the stealthiness parameter examined, *i.e.*,  $\chi$ (=0.45), the pair correlation function  $g_2(r)$  also possesses a sharp peak at r = D but does not exhibit either a power-law divergence for near contacts nor split second peaks; see Fig. 7. Furthermore, the small-scale oscillations in  $g_2(r)$  for  $\chi = 0.45$  for large r decay slightly faster than those with  $\chi = 0^+$ . Such differences in  $g_2(r)$  for the SHU ground states with  $\chi = 0^+$  and 0.45 arise because those with  $\chi = 0.45$  form only a small number of dimers but lack additional short-range order.

### 4. Conclusions and discussion

In sum, we have shown that one can use the modified collectivecoordinate optimization procedure [cf. eqn (2)] to create disordered SHU ground-state packings with packing fractions that far exceed those produced in our previous work within the disordered  $\chi$  regime,<sup>36,69</sup> thus advancing our understanding of the nature of this highly degenerate ground-state manifold. We revealed that in the small- $\chi$  limit, the ground-state packings associated with eqn (2) are hyperuniform, effectively jammed, and configurationally very near MRJ sphere packings, as measured by the same values of the packing fraction and mean contact number per particle, similar rattler concentrations, as well as pair statistics, both S(k) and  $g_2(r)$ , that are strikingly virtually identical to one another. The extraordinary existence of nonstealthy hyperuniform MRJ-like packings in the degenerate SHU ground-state manifold is possible because the cardinality of this infinite set is maximized as the stealthiness parameter  $\chi$ tends to zero. On the other hand, while our ground-state packings have a Dirac-delta distribution in packing fraction for finite N, numerically created MRJ packings have a range of packing fractions for finite N, which becomes a Dirac-delta distribution only in the thermodynamic limit.

We also showed that the modified optimization technique yields a large family of SHU packings for all values of  $\gamma < 1/2$ that exhibit other novel structural features. The particles in these disordered packings always form interparticle contacts, albeit with sparser contact networks, and have heretofore unattained maximal packing fractions  $\phi_{max}(\chi)$  compared to previous works on stealthy hyperuniform systems. In the  $\chi$ - $\phi_{\rm max}$  plane, the regions below and above the function  $\phi_{\rm max}(\chi)$ are satisfiable and unsatisfiable phases, respectively; see Fig. 3(a). By increasing  $\chi$  from 0 to 1/2, one can tune the contact networks of the SHU ground states from those that percolate with various degrees of connectivity for small to intermediate  $\chi$ values to those that do not percolate at higher  $\chi$  values characterized by dimer polymer-like chains. Detailed analyses of local structural statistics of these 3D SHU packings as a function of system size for  $\chi > 0$  and their lower-dimensional counterparts are reported in ref. 59.

Paper

The capability to attain ultradense SHU packings for all  $\chi$  within the disordered regime opens up exploration of new applications in optics and acoustics using such exotic amorphous materials.<sup>33,35,36,38,42–44,47,69</sup> Indeed, Vanoni *et al.*<sup>77</sup> recently showed how certain dynamical physical properties (*i.e.*, effective dynamic dielectric constant and time-dependent diffusion spreadability) of two-phase media derived from them are improved for a range of  $\chi$  within the disordered regime and packing fractions  $\phi$ .

Finally, we note that it has been a great computational challenge to generate the ideal MRJ state, which must be free of rattlers (defects).<sup>78</sup> Rapid-compression algorithms have been used over the years to generate such packings,<sup>5,6,60,79</sup> but they produce a small but significant fraction of rattlers that prevent the ideal MRJ state from truly forming due to a critical slowing down.<sup>80</sup> In future work, it would be interesting to explore the use of our effectively jammed but hyperuniform packing states as improved initial conditions for the Torquato–Jiao linear programming packing algorithm, which is designed to find inherent structures efficiently,<sup>60</sup> to possibly reduce the fraction of rattlers below 1.5%, thereby approaching the ideal state.

## Author contributions

S. T.: conceptualization, investigation, data curation, writing, funding acquisition; J. K.: data curation, software, investigation, writing.

## Data availability

The data that support the findings of this study are available from the corresponding author upon reasonable request.

## Conflicts of interest

There are no conflicts to declare.

## **Appendices**

### A. Simulation parameters

Here, we tabulate the simulation parameters for 3D ultradense SHU packings employed in this work.

### B. Total potential in the limit of $\chi \to 0$

We provide an explicit expression of the total potential (2) with  $\chi = \chi_{\min}(N)$  of *N*-particle systems (denoted by  $\chi = 0^+$  in Section 3):

$$\Phi(\mathbf{r}^{N}) = \rho v_0[S(k_{\min}\hat{\mathbf{e}}_1) + S(k_{\min}\hat{\mathbf{e}}_2) + S(k_{\min}\hat{\mathbf{e}}_3)] + \sum_{i < j} u(r_{ij}),$$
(9)

where  $k_{\min} = 2\pi/L$  is the smallest wavenumber of the reciprocal lattice vectors for the periodic simulation box, and  $\hat{e}_i$  (i = 1, 2, 3) are the standard basis of  $\mathbb{R}^3$ . Importantly, we approach the limit of  $\chi \to 0$  by increasing *N* in eqn (9) as large as possible (*i.e.*, *N* = 10774), ensuring that the obtained ground states are

hyperuniform in this limit, which is challenging for conventional compression algorithms to achieve.

## Acknowledgements

We thank Peter Morse, Charles Maher, and Paul Steinhardt for their very helpful remarks. This research was sponsored by the Army Research Office, accomplished under Cooperative Agreement Number W911NF-22-2-0103 and the National Science Foundation under Award No. CBET-2133179.

## Notes and references

- 1 S. Torquato and F. H. Stillinger, *Phys. Rev. E*, 2003, 68, 041113.
- 2 S. Torquato, Phys. Rep., 2018, 745, 1-95.
- 3 F. J. Dyson, Commun. Math. Phys., 1970, 19, 235-250.
- 4 H. L. Montgomery, Am. Math. Soc., 1973, 181–193.
- 5 A. Donev, F. H. Stillinger and S. Torquato, *Phys. Rev. Lett.*, 2005, **95**, 090604.
- 6 C. E. Maher, Y. Jiao and S. Torquato, *Phys. Rev. E*, 2023, **108**, 064602.
- 7 Y. Jiao, T. Lau, H. Hatzikirou, M. Meyer-Hermann, J. C. Corbo and S. Torquato, *Phys. Rev. E*, 2014, **89**, 022721.
- 8 A. Mayer, V. Balasubramanian, T. Mora and A. M. Walczak, *Proc. Nat. Acad. Sci. U. S. A.*, 2015, **112**, 5950–5955.
- 9 S. Torquato, A. Scardicchio and C. E. Zachary, J. Stat. Mech.: Theory Exp., 2008, P11019.
- 10 H. Wang, R. Samajdar and S. Torquato, *Phys. Rev. B*, 2024, 110, 104201.
- 11 C. Reichhardt and C. J. O. Reichhardt, *Soft Matter*, 2014, **10**, 7502–7510.
- 12 D. Hexner and D. Levine, Phys. Rev. Lett., 2015, 114, 110602.
- 13 D. Hexner, P. M. Chaikin and D. Levine, *Proc. Nat. Acad. Sci.* U. S. A., 2017, **114**, 4294–4299.
- 14 H. Zhang, X. Wang, J. Zhang, H.-B. Yu and J. F. Douglas, *Eur. Phys. J. E: Soft Matter Biol. Phys.*, 2023, **46**, 50.
- 15 Q.-L. Lei and R. Ni, Proc. Nat. Acad. Sci. U. S. A., 2019, 116, 22983–22989.
- 16 C. E. Zachary and S. Torquato, J. Stat. Mech.: Theory Exp., 2009, P12015.
- 17 E. C. Oğuz, J. E. S. Socolar, P. J. Steinhardt and S. Torquato, *Phys. Rev. B*, 2017, **95**, 054119.
- 18 S. Torquato, G. Zhang and M. de Courcy-Ireland, *J. Phys. A: Math. Theor.*, 2019, **52**, 135002.
- 19 O. H. E. Philcox and S. Torquato, *Phys. Rev. X*, 2023, 13, 011038.
- 20 A. Chremos and J. F. Douglas, *Phys. Rev. Lett.*, 2018, 121, 258002.
- 21 D. Chen, R. Samajdar, Y. Jiao and S. Torquato, *Proc. Natl. Acad. Soc.*, 2025, **122**, e2416111122.
- 22 O. U. Uche, F. H. Stillinger and S. Torquato, *Phys. Rev. E*, 2004, **70**, 046122.
- 23 R. D. Batten, F. H. Stillinger and S. Torquato, *J. Appl. Phys.*, 2008, **104**, 033504.

- 24 S. Torquato, G. Zhang and F. H. Stillinger, *Phys. Rev. X*, 2015, 5, 021020.
- 25 G. Zhang, F. H. Stillinger and S. Torquato, *Soft Matter*, 2017, 13, 6197–6207.
- 26 S. Ghosh and J. L. Lebowitz, Commun. Math. Phys., 2018, 363, 97-110.
- 27 M. Salvalaglio, D. J. Skinner, J. Dunkel and A. Voigt, *Phys. Rev. Res.*, 2024, **6**, 023107.
- 28 M. Florescu, S. Torquato and P. J. Steinhardt, Proc. Natl. Acad. Sci. U. S. A., 2009, 106, 20658–20663.
- 29 G. Gkantzounis, T. Amoah and M. Florescu, *Phys. Rev. B*, 2017, **95**, 094120.
- 30 L. S. Froufe-Pérez, M. Engel, J. José Sáenz and F. Scheffold, Proc. Nat. Acad. Sci. U. S. A., 2017, 114, 9570–9574.
- M. Castro-Lopez, M. Gaio, S. Sellers, G. Gkantzounis, M. Florescu and R. Sapienza, *APL Photonics*, 2017, 2, 061302.
- 32 S. Torquato and D. Chen, *Multifunct. Mater.*, 2018, 1, 015001.
- 33 F. Bigourdan, R. Pierrat and R. Carminati, *Opt. Express*, 2019, 27, 8666–8682.
- 34 V. Romero-García, N. Lamothe, G. Theocharis, O. Richoux and L. M. García-Raffi, *Phys. Rev. Appl.*, 2019, **11**, 054076.
- 35 A. Rohfritsch, J.-M. Conoir, T. Valier-Brasier and R. Marchiano, *Phys. Rev. E*, 2020, **102**, 053001.
- 36 S. Torquato and J. Kim, Phys. Rev. X, 2021, 11, 021002.
- 37 O. Christogeorgos, H. Zhang, Q. Cheng and Y. Hao, *Phys. Rev. Appl.*, 2021, 15, 014062.
- 38 H. Zhang, Q. Cheng, H. Chu, O. Christogeorgos, W. Wu and Y. Hao, *Appl. Phys. Lett.*, 2021, **118**, 101601.
- 39 V. Romero-García, É. Chéron, S. Kuznetsova, J.-P. Groby, S. Félix, V. Pagneux and L. M. Garcia-Raffi, *APL Mater.*, 2021, 9, 101101.
- 40 M. A. Klatt, P. J. Steinhardt and S. Torquato, *Proc. Nat. Acad. Sci. U. S. A.*, 2022, **119**, e2213633119.
- 41 N. Tavakoli, R. Spalding, A. Lambertz, P. Koppejan, G. Gkantzounis, C. Wan, R. Rohrich, E. Kontoleta, A. F. Koenderink, R. Sapienza, M. Florescu and E. Alarcon-Llado, *ACS Photonics*, 2022, 9, 1206–1217.
- 42 K. Tang, Y. Wang, S. Wang, D. Gao, H. Li, X. Liang, P. Sebbah, Y. Li, J. Zhang and J. Shi, *Phys. Rev. Appl.*, 2023, **19**, 054035.
- 43 M. Merkel, M. Stappers, D. Ray, C. Denz and J. Imbrock, *Adv. Photonics Res.*, 2023, 5, 2300256.
- 44 L. Alhatz, J.-M. Conoir and T. Valier-Brasier, *Phys. Rev. E*, 2023, **108**, 065001.
- 45 N. Granchi, M. Lodde, K. Stokkereit, R. Spalding,
  P. J. Veldhoven, R. Sapienza, A. Fiore, M. Gurioli,
  M. Florescu and F. Intonti, *Phys. Rev. B*, 2023, **107**, 064204.
- 46 H. Zhuang, D. Chen, L. Liu, D. Keeney, G. Zhang and Y. Jiao, J. Phys.: Condens. Matter, 2024, 36, 285703.
- 47 S. Kuznetsova, J. P. Groby, L. M. Garcia-Raffi and V. Romero-García, *Waves Random Complex Media*, 2024, **34**, 1878–1896.
- 48 E. Wigner, Phys. Rev., 1934, 46, 1002.
- 49 R. P. Feynman and M. Cohen, *Phys. Rev.*, 1956, **102**, 1189–1204.
- 50 R. B. Laughlin, *The Quantum Hall Effect*, Springer, 1987, pp. 233–301.

- 51 C. Broholm, R. J. Cava, S. A. Kivelson, D. G. Nocera, M. R. Norman and T. Senthil, *Science*, 2020, 367, eaay0668.
- 52 G. Semeghini, H. Levine, A. Keesling, S. Ebadi, T. T. Wang, D. Bluvstein, R. Verresen, H. Pichler, M. Kalinowski, R. Samajdar, A. Omran, S. Sachdev, A. Vishwanath, M. Greiner, V. Vuletić and M. D. Lukin, *Science*, 2021, 374, 1242–1247.
- 53 R. Samajdar, W. W. Ho, H. Pichler, M. D. Lukin and S. Sachdev, *Proc. Natl. Acad. Soc.*, 2021, **118**, e2015785118.
- 54 S. Kivelson and S. Sondhi, Nat. Rev. Phys., 2023, 5, 368-369.
- 55 G. Zhang, F. Stillinger and S. Torquato, *Phys. Rev. E*, 2015, **92**, 022119.
- 56 P. K. Morse, J. Kim, P. J. Steinhardt and S. Torquato, *Phys. Rev. Res.*, 2023, **5**, 033190.
- 57 A. Shih, M. Casiulis and S. Martiniani, *Phys. Rev. E*, 2024, **110**, 034122.
- 58 G. Zhang, F. H. Stillinger and S. Torquato, *J. Chem. Phys.*, 2016, **145**, 244109.
- 59 J. Kim and S. Torquato, J. Chem. Phys., 2025, in press; arXiv, 2025, preprint, arXiv:2504.16924, https://arxiv.org/abs/2504. 16924.
- 60 S. Torquato and Y. Jiao, Phys. Rev. E, 2010, 82, 061302.
- 61 P. Charbonneau, E. I. Corwin, G. Parisi and F. Zamponi, *Phys. Rev. Lett.*, 2012, **109**, 205501.
- 62 F. H. Stillinger, E. A. DiMarzio and R. L. Kornegay, *J. Chem. Phys.*, 1964, **40**, 1564–1576.
- 63 C. S. O'Hern, S. A. Langer, A. J. Liu and S. R. Nagel, *Phys. Rev. Lett.*, 2002, 88, 075507.
- 64 S. Wilken, R. E. Guerra, D. Levine and P. M. Chaikin, *Phys. Rev. Lett.*, 2021, **127**, 038002.
- 65 S. Atkinson, F. H. Stillinger and S. Torquato, *Phys. Rev. E*, 2013, **88**, 062208.
- 66 A. Donev, S. Torquato and F. H. Stillinger, *Phys. Rev. E*, 2005, 71(011105), 1–14.
- 67 S. Torquato, Int. J. Solids Structures, 2000, 37, 411-422.
- 68 S. Torquato and F. H. Stillinger, Rev. Mod. Phys., 2010, 82, 2633.
- 69 J. Kim and S. Torquato, Proc. Natl. Acad. Sci. U. S. A., 2020, 117, 8764–8774.
- 70 G. Matheron, Cahiers de Géostatistique, 1993, 107, 107-113.
- 71 S. Franz, G. Parisi, M. Sevelev, P. Urbani and F. Zamponi, *Sci. Phys.*, 2017, **2**, 019.
- 72 C. S. O'Hern, L. E. Silbert, A. J. Liu and S. R. Nagel, *Phys. Rev. E*, 2003, 68, 011306.
- 73 Y. Jin and H. Yoshino, Proc. Natl. Acad. Sci. U. S. A., 2021, 118, e2021794118.
- 74 J. Nocedal, Math. Comput., 1980, 35, 773-782.
- 75 D. C. Liu and J. Nocedal, Math. Program., 1989, 45, 503-528.
- 76 M. E. J. Newman and R. M. Ziff, Phys. Rev. Lett., 2000, 85, 4104–4107.
- 77 C. Vanoni, J. Kim, P. J. Steinhardt and S. Torquato, Dynamical properties of particulate composites derived from ultradense stealthy hyperuniform sphere packings, 2025.
- 78 S. Torquato, Phys. Rev. E, 2021, 103, 052126.
- 79 S. Torquato, T. M. Truskett and P. G. Debenedetti, *Phys. Rev. Lett.*, 2000, 84, 2064–2067.
- 80 S. Atkinson, G. Zhang, A. B. Hopkins and S. Torquato, *Phys. Rev. E*, 2016, **94**, 012902.