Hyperuniform jammed sphere packings have anomalous material properties

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A spatial distribution is hyperuniform if it has local density fluctuations that vanish in the limit of long length scales. Hyperuniformity is a well known property of both crystals and quasicrystals. Of recent interest, however, is disordered hyperuniformity; the presence of hyperuniform scaling without long-range configurational order. Jammed granular packings have been proposed as an example of disordered hyperuniformity, but recent numerical investigation has revealed that many jammed systems instead exhibit a complex set of distinct behaviors at long, emergent length scales. We use the Voronoi tessellation as a tool to define a set of rescaling transformations that can impose hyperuniformity on an arbitrary weighted point process, and show that these transformations can be used in simulations to iteratively generate hyperuniform, mechanically stable packings of athermal soft spheres. These hyperuniform jammed packings display atypical mechanical properties, particularly in the low-frequency phononic excitations, which exhibit an isolated band of highly collective modes and a band gap around zero frequency.

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I. INTRODUCTION

A statistically homogeneous distribution of points throughout some volume is uniform in the sense that the probability density function is constant throughout the volume. However, any finite sampling of this function will necessarily contain density fluctuations at all length scales. By contrast, a hyperuniform random distribution is one in which density fluctuations are completely suppressed at long length scales [1,2]. While this phenomenon is most closely associated with crystalline and quasicrystalline patterns [3], hyperuniformity also appears in disordered systems such as the prime numbers [4,5] as well as in materials which are both disordered and statistically isotropic [3,6–13]. Hyperuniformity can even be realized in amorphous packings of circles and spheres [14–18]. Further, hyperuniform materials could be used for creating efficient sensory networks without the drawbacks of aliasing. These hyperuniform sensory networks have even been shown to arise in nature, in particular in the distribution of photocells within the eyes of chickens [19].

It is natural to suspect that all jammed sphere packings are hyperuniform as the relaxation process by which such packings are formed will necessarily involve smoothing out local density fluctuations relative to an uncorrelated initial configuration [1]. However, recent numerical evidence demonstrates that, for typical protocols used to create jammed packings in silico, an emergent length scale appears at which this suppression abates [15,20,21]. Further exploration of the link between hyperuniformity and jamming could give rise to deeper insights into the nature of the glass transition, and more generally of geometric order within amorphous materials. In this light, any protocol which can generate concrete examples of hyperuniform materials has significant theoretical and numerical applications.

II. CREATING HYPERUNIFORM PACKINGS

In order to create hyperuniform jammed packings, we combine a protocol for creating mechanically stable sphere packings with a straightforward protocol for transforming any weighted point pattern into a hyperuniform point pattern. The mechanism for transforming point patterns into hyperuniform point patterns bears strong similarities with Ref. [17], which proved that such point patterns will be type I hyperuniform. This algorithm was developed in parallel with Ref. [18], which demonstrated that thermal packings created in this way are unusually stable. Just as in those works, our methods involve an iterative process of rescaling individual particles to a uniform local volume fraction within each Voronoi cell. In Ref. [17] this process is applied once and the resulting hyperuniform weighted point patterns are studied. In Ref. [18] the resulting packing is then relaxed and the process is repeated until the standard deviation of the local volume fraction distribution no longer changes. By contrast, in our work we find a broad range of phase space above jamming in which we can continue this process until the local volume fraction fluctuations are brought identically to zero. Further, the area of applicability is significantly different in that our work focuses on the overjammed regime of soft-particle packings, which allows us to directly explore the implications on mechanical properties for exactly hyperuniform sphere packings. As a technical advancement, in this work we introduce a process to simultaneously perform the relaxation (energy minimization) and the iterative rescaling of radii, resulting in a significant increase in speed and efficiency of this algorithm.

A. Mechanically stable sphere packings

Packings are simulated as sets of $N$ spherical particles each in a cubic periodic bounding box of unit volume. Particles have radii $r_i$ and positions $\mathbf{x}_i$, and interact through a harmonic
FIG. 1. The cells fully (light orange) and partially (orange) within a circular region. (a) In a periodic tessellation, which is hyperuniform, the variance within a sampling window is introduced only from the cells on the window’s boundary. (b) In the tessellation of a Poisson point pattern, variance arises from variation in both the bulk and the surface of the window, leading to uniform statistics. If, however, the weight of each cell is made to be proportional to its bulk and the surface of the window, leading to uniform statistics. If, however, the weight of each cell is made to be proportional to its area, then there will be no density fluctuations in the interior and as such it will follow hyperuniform statistics.

pairwise interaction:

$$U = \sum_{i<j} \frac{1}{2} \left( 1 - \frac{|\vec{x}_j - \vec{x}_i|}{r_i + r_j} \right)^2 \Theta \left( 1 - \frac{|\vec{x}_j - \vec{x}_i|}{r_i + r_j} \right).$$  \hspace{1cm} (1)$$

where $\Theta$ is the Heaviside step function.

Packings are created from random starting positions at a fixed packing fraction $\varphi$ and then energy-minimized using the FIRE algorithm until the packing reaches mechanical stability [22]. These relaxation methods are chosen to approximate a rapid quench from infinite temperature [23].

B. From weighted point pattern to hyperuniformity

Given a point pattern in $d$ dimensions, we may describe density fluctuations by considering the number of particles which fall within an ensemble of similarly shaped window functions. We define $\sigma^2_r(r)$ to be the variance of the total number to fall within the subset of windows sharing a length scale $r$.

As illustrated in Fig. 1, a Poisson point pattern, or indeed any sufficiently uncorrelated distribution, will have a variance which grows linearly with the mean number of points and so will have $\sigma^2_r(r) \propto r^d$. Conversely, in any periodic or quasicrystalline distribution, the only contribution to variance within a sampling window comes from those unit cells which are bisected by the window’s boundary, leading to a variance which grows as the surface area $\sigma^2_r(r) \propto r^{d-1}$. Such a point pattern is said to be hyperuniform. Within the context of jammed packings, it is convenient to measure $\varphi_i$, the local volume fraction within a region, instead of the number of particles within a region. As $\varphi_i$ is proportional to the number of particles divided by the volume of a region, this hyperuniform condition may be recast in terms of the local packing fraction by division by $r^{2d}$. Thus, we find that the variance of local packing fractions for normally distributed points scales as $r^{-d}$ and the hyperuniform condition is now

$$\langle \varphi_i^2 \rangle - \langle \varphi_i \rangle^2 \propto r^{-(d+1)}. \hspace{1cm} (2)$$

This definition may be generalized to weighted point patterns by considering the variance of the total weight within the window function rather than the number of points. For purposes of analyzing density fluctuations in granular packings, we idealize each particle as a Dirac delta distribution at the particle’s center with weight equal to the volume of the particle. This choice has been shown to not affect the long-range correlations [15]. Thus we compute the density distribution $\rho$ as

$$\rho(\vec{x}) = \sum_i \frac{\pi^{d/2}}{\Gamma \left( \frac{d}{2} + 1 \right)} r_i^{d/2} \delta(\vec{x} - \vec{x}_i). \hspace{1cm} (3)$$

We propose a method to create hyperuniform packings from conventional packings by iteratively resizing particles to achieve uniform local density in the system, as illustrated in Fig. 2. Similarly to work done by Morse et al. [24], we exploit the geometric cell properties of the Voronoi tessellation in a minimization scheme. We first turn the point pattern associated with the packing into a hyperuniform point pattern with the following algorithm:

2. Compute the volume of each cell in the tessellation.
3. Assign to each point a new weight proportional to the volume of the local cell.

In Fig. 3, we demonstrate the validity of this method for creating hyperuniform distributions. We do so by applying this Voronoi rescaling to a large number of Poisson points in a periodic box. We note that although we use the Voronoi tessellation, any tessellation with a well-behaved diameter distribution should work equally well.

In the context of sphere packings, this process will change each particle’s volume and thus the polydispersity of the...
III. ANALYSIS

A. Convergence

The dynamics of our iterative algorithm are not monotonic in energy, and explore a particularly high-dimensional configuration space of positions and radii in which we expect hyperuniform fixed points to be relatively rare. It is thus surprising that this algorithm would ever converge, much less as robustly as it does.

We track the dynamics of iteration by measuring the maximum displacement, which provides an indication of this convergence. Sufficiently far from the jamming point, we find the behavior shown in Fig. 4: systems quickly enter a regime of random motion within a consistent range of maximum displacements. Eventually, the system (usually) falls into an approximately exponential descent until the displacement becomes impossible to resolve numerically. This reduction in displacement is connected to the existence of a fixed point, and the exponential behavior is related to the local analyticity of the energy landscape about that fixed point. Some systems instead remain in a state of random motion for a very long time, and limit cycles, while rare, have been observed.

Over an ensemble of many similar realizations, we observe that the probability of convergence approaches its maximum exponentially. This can be understood by assuming that falling into a fixed point is a Poisson process on the random walk through configuration space. The long-time probability of convergence depends on the parameters of the realizations, and most sensitively on the packing fraction, which we illustrate in Fig. 5. While the characteristic time of convergence increases mildly as the jamming transition \( \varphi_j \) is approached, the total probability of convergence decreases, carving out a regime around the critical point in which locating fixed points is at best computationally infeasible and at worst impossible. Larger systems show qualitatively similar behavior; however we have not systematically studied the finite-size scaling of this convergence when close to the jamming transition as the timescales involved rapidly diverge for larger \( N \). In order to ensure relatively quick convergence to fixed points, we primarily focus on systems relatively far from jamming.
Qualitatively similar behavior. Larger systems show convergence of packings with density $\phi_s$ at $\varphi \approx 0.64$. Packings nearer to jamming converge less frequently, and those that do converge do so more slowly.

The static structure factor for weighted points measures the strength of density fluctuations at a given wave vector $\vec{k}$ and is defined as

$$S(\vec{k}) = \frac{1}{N} \rho(\vec{k}) \rho(-\vec{k}).$$

Any point pattern will have large fluctuations at short distances (large $\vec{k}$). Uniform point patterns will have nonzero fluctuations even at very large distances ($\vec{k}$ approaching zero) whereas a hyperuniform point pattern will have vanishing fluctuations [1].

In Fig. 6, we show that iteration of our constructive algorithm suppresses long-range density fluctuations, as measured by the average value of $S(\vec{k})$ for a set of packings. Of particular interest is $\lim_{|\vec{k}| \rightarrow 0} S(\vec{k})$ which must be zero when we reach a hyperuniform fixed point. In the initial mechanically stable packing we observe that $S(\vec{k})$ reaches a constant value as $\vec{k}$ approaches zero, indicating that it is not hyperuniform. We consider the value of $S(\vec{k})$ at $\vec{k}_{\text{min}}$, where $\vec{k}_{\text{min}}$ is the wave number for which $S(\vec{k})$ takes its minimum value. After a single iteration this value of $S(\vec{k}_{\text{min}})$ increases, because individual iterations only perform partial minimizations, which initially introduces density fluctuations. As iteration continues, however, $S(\vec{k}_{\text{min}})$ decreases, indicating that the system is becoming hyperuniform. Near the fixed point, the functional form of the structure factor resembles a power law $S(\vec{k}) \approx a|\vec{k}|^{1/2}$. That this power law appears to be nonanalytical as $\vec{k}$ goes to 0 is surprising in light of the results of Ref. [17] which show that hyperuniform point patterns derived from Voronoi tesselations will have small-$\vec{k}$ behavior that scales as either $\vec{k}^2$ or $\vec{k}^4$. This discrepancy may arise from the fact that points in that work were placed at the centroids of the Voronoi cells whereas our particles are not constrained in that way. It is also possible that $S(\vec{k})$ continues to steepen at smaller $\vec{k}$ and thus approaches an even power law. Nevertheless, this difference is worthy of further study.

C. Mechanical properties

Packings generated by Voronoi iteration represent a concrete set of examples with which to explore the mechanical properties of hyperuniform granular materials. The vibrational density of states $D(\omega)$ for sphere packings above jamming has been well characterized in previous work [27–30]. In Fig. 7(a), we show the density of states for both the initial packing and the hyperuniform final configuration. At the densities explored in this work we observe, as expected, a low-frequency plateau associated with the boson peak and then an $\omega^2$ falloff at the lowest frequencies in our initial packings. The final hyperuniform configurations, however, have a remarkably different density of states: the lowest frequency scaling regime is entirely absent, replaced by an abrupt low-frequency cutoff at a finite frequency forming a phononic band gap about zero frequency. Above this gap, there is a discrete band of modes which we refer to as “sloshing modes.” The high-frequency behavior, however, remains largely unchanged.

Figure 7(b) shows the inverse participation ratio (IPR) as a function of frequency, defined for a given eigenvector as

$$\text{IPR} = \frac{\sum_i \|\vec{v}_i\|^4}{\left(\sum_i \|\vec{v}_i\|^2\right)^2}.$$
where $\vec{v}_i$ is the mode displacement of the $i$th particle. This quantity measures the localization of each mode. The modes that constitute the sloshing band are easily distinguished from the initial configurations as they are highly collective, with typical IPR lower than at any other point in the spectrum. The IPR for frequencies above the sloshing band is quantitatively similar to that observed in the initial configurations.

In addition to being collective, these modes are highly spatially correlated, consisting of nearly affine “sloshing” motions over large portions of the system, visualized in Fig. 8. To quantify this, we define the local affinity (LA) of a single mode $\vec{v}_i$ as

$$
\text{LA} = \frac{1}{N_{\text{neigh}}} \sum_{i,j} \frac{\vec{v}_i \cdot \vec{v}_j}{\|\vec{v}_i\| \|\vec{v}_j\|}.
$$

where $i$ indicates the particle index and $(i, j)$ denotes the $N_{\text{neigh}}$ neighbors in the radical Delaunay triangulation [25]. For each mode this quantity measures the extent to which adjacent particles oscillate in phase and in the same direction, having a value of 1 for a spatially affine displacement of the entire system. Figure 7(c) shows that when we measure the local affinity, we again find a clear distinction between the sloshing modes in the hyperuniform systems and the much less correlated low-frequency modes of the initial configurations. Modes within the sloshing band are consistently more affine than the rest.

IV. CONCLUSIONS

We have demonstrated that hyperuniform, overjammed, polydisperse granular packings not only exist, but can be efficiently generated at large system size through a process of iterative optimization. This result allows us to probe a concrete set of systems to better understand the implications of hyperuniformity.

Despite being chosen on geometrical grounds, the hyperuniform packings we construct exhibit unusual mechanical properties. We have investigated these by exploring their linear vibrational modes. Here we find a low-energy band gap structure not seen in typical preparations, which includes highly collective excitations and regions of reduced density of states, indicative of the formation of phononic band gaps. Further modification of the band structure of granular media may be possible through similar geometric optimization.

Additionally, the lack of normal modes at low frequency suggests that these packings are highly stable against mechanical perturbation. Accordingly we believe that they lie in particularly deep wells in the soft sphere energy landscape. This suggests that the procedure of smoothing out density fluctuations through Voronoi iteration may be somewhat analogous to aging in thermal glasses. Further elaboration of this correspondence could shed light on the geometric properties of aged glasses, and the role of large-scale geometric heterogeneity in glassy dynamics.
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