Numerical test of the Edwards conjecture shows that all packings are equally probable at jamming

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In the late 1980s, Sam Edwards proposed a possible statistical-mechanical framework to describe the properties of disordered granular materials1. A key assumption underlying the theory was that all jammed packings are equally likely. In the intervening years it has never been possible to test this bold hypothesis directly. Here we present simulations that provide direct evidence that at the unjamming point, all packings of soft repulsive particles are equally likely, even though generically, jammed packings are not. Typically, jammed granular systems are observed precisely at the unjamming point since grains are not very compressible. Our results therefore support Edwards’ original conjecture. We also present evidence that at unjamming the configurational entropy of the system is maximal.

In science, most breakthroughs cannot be derived from known physical laws: they are based on inspired conjectures2. Comparison with experiment of the predictions based on such a hypothesis allows us to eliminate conjectures that are clearly wrong. However, there is a distinction between testing the consequences of a conjecture and testing the conjecture itself. A case in point is Edwards’ theory of granular media. In the late 1980s, Edwards and Oakeshott1 proposed that many of the physical properties of granular materials (‘powders’) could be predicted using a theoretical framework that was based on the assumption that all distinct packings of such a material are equally likely to be observed. The logarithm of the number of such packings was postulated to play the same role as entropy does in Gibbs’ statistical-mechanical description of the thermodynamic properties of equilibrium systems. However, statistical-mechanical entropy and granular entropy are very different objects. Until now, the validity of Edwards’ hypothesis could not be tested directly—mainly because the number of packings involved is so large that direct enumeration is utterly infeasible—and, as a consequence, the debate about the Edwards hypothesis has focused on its consequences, rather than on its assumptions. Here we present results that show that now, at last, it is possible to test Edwards’ hypothesis directly by numerical simulation. To our own surprise, we find that the hypothesis appears to be correct precisely at the point where a powder is just at the (un)jamming threshold. However, at higher densities, the hypothesis fails. At the unjamming transition, the configurational entropy of jammed states appears to be at a maximum.

The concept of ‘ensembles’ plays a key role in equilibrium statistical mechanics, as developed by J. Willard Gibbs, well over a century ago3. The crucial assumption that Gibbs made to arrive at a tractable theoretical framework to describe the equilibrium properties of gases, liquid and solids was that, at a fixed total energy, every state of the system is equally likely to be observed. The distinction between, say, a liquid at thermal equilibrium and a granular material is that in a liquid, atoms undergo thermal motion whereas in a granular medium (in the absence of outside perturbations) the system is trapped in one of (very) many local potential energy minima. Gibbsian statistical mechanics cannot be used to describe such a system. The great insight of Edwards was to propose that the collection of all stable packings of a fixed number of particles in a fixed volume might also play the role of an ‘ensemble’ and that a statistical mechanics-like formalism would result if one assumed that all such packings were equally likely to be observed, once the system had settled into a mechanically stable ‘jammed’ state. The nature of this ensemble has been the focus of many studies4–6.

Jamming is ubiquitous and occurs in materials of practical importance, such as foams, colloids and grains when they solidify in the absence of thermal fluctuations. Decompressing such a solid to the point where it can no longer achieve mechanical equilibrium leads to unjamming. Studies of the unjamming transition in systems of particles interacting via soft, repulsive potentials have shown that this transition is characterized by power-law scaling of many physical properties7–11. However, the exact nature of both the ensemble of jammed states and the unjamming transition remains unclear.

Here, we report a direct test of the Edwards conjecture, using a numerical scheme for computing basin volumes of distinct jammed states (energy minima) of N polydisperse, frictionless discs held at a constant packing fraction φ. Uniquely, our numerical scheme allows us to compute Ω, the number of distinct jammed states, and the individual probabilities $p_i \propto 1/\ln \Omega$ of each observed packing to occur. Figure 1a shows a snapshot of a section of the system, consisting of particles with a hard core and a soft shell. We obtain jammed packings by equilibrating a hard-disc fluid and inflicting the particles instantaneously to obtain the desired soft-disc volume fractions (φ), followed by energy minimization (see Methods). The minimization procedure finds individual stable packings with a probability $p_i \propto 1/\ln \Omega$ of each observed packing to occur. Figure 1a shows a snapshot of a section of the system, consisting of particles with a hard core and a soft shell. We obtain jammed packings by equilibrating a hard-disc fluid and inflicting the particles instantaneously to obtain the desired soft-disc volume fractions (φ), followed by energy minimization (see Methods). The minimization procedure finds individual stable packings with a probability $p_i \propto 1/\ln \Omega$ of each observed packing to occur. Averages computed using this procedure, represented by $\langle \cdots \rangle_\Omega$, would then lead to a bias originating from the different $\nu_i$ values. Recent advances in numerical methods12–14 have now enabled direct computation of $\nu_i$ and therefore, an unbiased characterization of the phase space.

We report a detailed analysis of the distribution of $\nu_i$ for a fixed number of discs $N = 64$ (all maximum system sizes in our study were set by the current limits on computing power). We compute $\nu_i$ using a thermodynamic integration scheme13,14, and compute the average basin volume $\langle \nu \rangle(\phi)$. The number of jammed states is, explicitly, $\Omega(\phi) = V(\phi)/\langle \nu \rangle(\phi)$, where $V(\phi)$ is the total available phase-space volume at a given φ. A convenient way to check equiprobability is to compare the Boltzmann entropy $S_B = \ln \Omega - \ln N!$, which counts all packings with the same weight, and the Gibbs entropy $S_G = -\sum_i p_i \ln p_i - \ln N!$ (refs 17–19). The Gibbs entropy satisfies $S_G \leq S_B$, saturating the bound when all $p_i$ are equal: $p_i(1/\ln \Omega) = 1/\Omega$. As shown in Fig. 2a, $S_G$ approaches

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Figure 1 | Snapshot of the system studied and illustration of the associated energy landscape at different volume fractions. a, Snapshot of a jammed packing of discs with a hard core (dark shaded regions) plus soft repulsive corona (light shaded regions). b,c, Illustration of configurational space for jammed packings. The hatched regions are inaccessible due to hard-core overlaps. Single-coloured regions with contour lines represent the basins of attraction of distinct minima. The dark blue region with solid dots indicates the coexisting unjammed fluid region and hypothetical marginally stable packings, respectively. The volume occupied by the fluid $V_{\text{unj}}$ is significant only for finite-size systems at or near unjamming. When $\phi \gg \phi^*$ (b) the distribution of basin volumes is broad but as $\phi \to \phi^*$ (c) the distribution of basin volumes approaches a delta function satisfying Edwards’ hypothesis.

Figure 2 | Numerical results obtained by basin-volume calculations for jammed packings of $N = 64$ discs with a hard core and a soft shell. a, Gibbs entropy $S_G$ and Boltzmann entropy $S_B$ as a function of volume fraction. $S_B$ is computed both parametrically by fitting $B(f)$ with a generalized Gaussian function (Gauss) and non-parametrically by computing a kernel density estimate (KDE) as in ref. 13. The dashed curves are a second-order polynomial fit. b, Scatter plot of the negative log-probability of observing a packing, $-\ln p_i = f_i + \ln V_i(\phi)$, where $V_i$ is the accessible fraction of phase space (see Methods) as a function of log-pressure, $A$. The black solid lines are lines of best fit computed by linear minimum mean square error using a robust covariance estimator and bootstrap (see Methods). c,d, Slopes $\lambda(\phi)$ (c) and intercepts $c(\phi)$ (d) of linear fits for equation (16). Solid lines are lines of best fit and error bars refer to the standard error computed by bootstrap.

$S_B$ from below as $\phi \to \phi^{(c)}_{\text{min}} \approx 0.82$. Fig. 1b,c schematically illustrates the evolution of the basin volumes as the packing fraction is reduced.

To characterize the distribution of basin volumes, we analyse the statistics of $v_i$ along with the pressure $P_i$ of each packing. It is convenient to study $F_i \equiv -\ln v_i$ as a function of $A_i \equiv \ln P_i$. As shown in Fig. 2b, we observe a strong correlation between $F_i$ and $A_i$, which we quantify by fitting the data to a bivariate Gaussian distribution. The conditional expectation of $F$ given $A$ then yields a linear relationship (denoted by solid lines in Fig. 2b) such that $\langle F \rangle_B(\phi; A) \propto \lambda(\phi) A + c(\phi)$, where $\langle F \rangle_B(\phi; A)$ represents the average over all basins at a given $A$. Previous studies at higher packing fractions\(^{13}\) indicate that this relationship is preserved in the thermodynamic limit. Defining $f = F/N$, we have (see Methods for details)

$$\langle f \rangle_B(\phi; A) = \lambda(\phi) A + c(\phi)$$

$$= \lambda(\phi) A + f_B(\phi)$$

(1)
where $\Delta \Lambda = \Lambda - \langle \Lambda \rangle_B (\phi)$. For Edwards’ hypothesis to be valid, we require that in the thermodynamic limit: the distribution of volumes approaches a Dirac delta, which follows immediately from the fact that the variance $\sigma_1^2 \sim 1/N$ (ref. 15) (see Supplementary Information); and $F_i$ needs to be independent of $\Lambda$, as well as of all other structural observables (see Supplementary Information), and therefore $\langle\Lambda\rangle_B$ must necessarily vanish. As can be seen from Fig. 2c,d, within the range of volume fractions studied, $\lambda(\phi)$ decreases but saturates to a minimum as $\phi \to \phi_{s=\infty}$. We argue below that the saturation is a finite-size effect. An extrapolation using the linear regime in Fig. 2c indicates that $\lambda \to 0$ at packing fraction $\phi_{s=\infty} = 0.82 \pm 0.07$, remarkably close to where our extrapolation yields $S_c = S_0$. The analysis of basin volumes, therefore, strongly suggests that equiprobability is approached only at a characteristic packing fraction and that the vanishing of $\lambda(\phi)$ can be used to estimate the point of equiprobability.

We next show that $\lambda(\phi)$ does indeed tend to zero in the thermodynamic limit. We use the fluctuations $\sigma_2^2$, $\sigma_2^2$, and the covariance $\sigma_2^2$ of the elements of the covariance matrix $\hat{\sigma} = ([\sigma_{ij}, \sigma_{ij}], \sigma_{ij}, \sigma_{ij})$ of the joint distribution of $f$ and $\Lambda$ (see Methods for details), to define $\lambda$ and $c$ as:

$$
\lambda(\phi) \equiv \frac{\sigma_2^2(\phi)}{\sigma_2^2(\phi)} - \frac{\sigma_2^2(\phi)}{\sigma_2^2(\phi)} \langle \Lambda \rangle_B (\phi)
$$

From Fig. 2b we observe that the decrease of $\lambda$ is driven by $\sigma_2^2$ increasing to a maximum, while $\sigma_2^2$ and $\sigma_2^2$ decrease (see Supplementary Fig. 2). We expect the main features of these distributions to be preserved as the system size $N$ is increased, which suggests that for larger $N$, where basin volume calculations are still intractable for multiple densities, the maximum in $\sigma_2^2$ can be used to identify $\phi_{s=\infty}$. We have directly measured $\chi_N = N\sigma_2^2$ using our sampling scheme—which samples packings with probability proportional to the volume of their basin of attraction—for systems of up to $N = 128$ discs (see inset of Fig. 3a) and finite-size scaling indicates that $\chi_N$ diverges as $\phi \to \phi_{s=\infty} = 0.841(3)$ (see Supplementary Information). The saturation of $\lambda$ to a minimum as $\phi \to \phi_{s=\infty}$, for small $N$, is determined by the fact that $\chi_N$ diverges only in the thermodynamic limit; a detailed discussion is given in the Methods.

Interestingly, we find evidence that in the thermodynamic limit, the point of equiprobability, $\phi_{s=\infty}$, coincides with the point at which the system unjams, $\phi_{s=\infty}$. We use two characteristics of the unjamming transition to locate $\phi_{s=\infty}$: the average pressure of the packings goes to zero, and therefore $\langle \Lambda \rangle_B \to -\infty$ (see Fig. 3b); and the probability of finding jammed packings, $p_j$, goes to zero (see inset of Fig. 3b). A scaling analysis indicates that $\langle \Lambda \rangle_B \to -\infty$ as $\phi_{s=\infty} = 0.841(3)$, and $p_j \to 0$ as $\phi_{s=\infty} = 0.844(2)$ (see Supplementary Information). We thus find that $\phi_{s=\infty} = \phi_{s=\infty}$, within numerical error and up to corrections to finite-size scaling. Our simulations therefore lead to the surprising conclusion that the Edwards conjecture appears to hold precisely at the (un)jamming transition. We note that our earlier simulations, which were performed at densities much above jamming, did not support the equiprobability hypothesis. The earlier simulations were in fact far enough away from unjamming that the emergence of equiprobability at this point could not be anticipated. Our earlier findings, therefore, do not contradict the more recent ones and are completely consistent with these.

Why is $\chi_N$ related to the unjamming transition? As the particles interact via purely repulsive potentials, the pressure $P$ is strictly positive, which implies that the fluctuations of $P$ have a floor and go to zero at unjamming. The relative fluctuations $\chi_P \equiv N\sigma^2(P/(P)_B)$ can be non-zero, and a diverging $\chi_P$ would then imply a diverging $\chi_N$. Because of the bounded nature of $P$ (refs 21–23), however, $\chi_P$ can diverge only at the unjamming transition where $\langle P \rangle_B \to 0$ (see Methods). We find that $\chi_P$ does diverge (Fig. 3a) and finite-size scaling yields $\phi_{s=\infty} = 0.841(3)$, in agreement with what has been found for $\chi_A$. Returning to the $N = 64$ case that we have analysed using the basin volume statistics, we find that both $\chi_P$ and $\chi_A$ saturate to their maximum values over similar ranges of $\phi$ and our estimate $\phi_{s=\infty} \approx 0.82$, where $S_c = S_0$ and $\lambda \to 0$, falls in this region. In addition, the average number of contacts $\langle z_P(\phi_{s=\infty}) \rangle = 4.1 \pm 0.2$ is close to the isostatic value $z_{iso}^{(N=64)} = 2d - 2/64 \approx 3.97$ (ref. 10) (see Supplementary Information).

Finally, we note that the states in the generalized Edwards ensemble $^{5,21,22,23}$ characterized by $\phi$ and $P$ have basin volumes that are similar, if not identical, over the full range of $\phi$ that we have explored (see scatter plot in Fig. 2b), indicating that equiprobability in the stress–volume ensemble $^{5,24}$ is a more robust formulation.
of the Edwards hypothesis. This observation is consistent with recent experiments\(^{26}\).

Although, the equiprobability of jammed states at a given packing fraction was posited by Edwards for jammed packings of hard particles, our analysis shows that for soft particles, the Edwards hypothesis is valid only for the marginally jammed states at \(N_e \to \infty \), where the jamming probability vanishes, the entropy is maximized, and relative pressure fluctuations diverge. We have shown not only that there exists a practical ‘Edwardsian’ packing generation protocol, capable of sampling jammed states equiprobably, but we have uncovered an unexpected property of the energy landscape for this class of systems. At this stage, we cannot establish whether the same considerations are valid in three dimensions, although the already proven validity of equation (16) in three dimensions would suggest so\(^2\). The exact value of the entropy at unjamming, whether finite or not, also needs to be elucidated. The implications for ‘soft’ structural glasses is apparent: at \(N \to \infty \) the uniform size of the basins implies that the system, when thermalized, has the same probability of visiting all of its basins of attraction; hence, there are no preferred inherent structures. This could be a signature of the hard-sphere transition occurring at the same point\(^{12}\). Our approach can, therefore, be extended to spin-glasses and related problems, and it would be clearly very exciting to explore the analogies and differences between ‘jamming’ in various systems for which the configuration space can break up into many distinct basins.

**Methods**

Methods, including statements of data availability and any associated accession codes and references, are available in the online version of this paper.

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**Author contributions**

S.M., D.F. and B.C. designed the study. S.M. and K.J.S. developed the software and performed the numerical simulations. S.M., K.R., B.C. and D.F. analysed the data and wrote the manuscript.

**Additional information**

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**Competing financial interests**

The authors declare no competing financial interests.
Methods

Packing preparation protocol. In this section we describe the algorithm that we have used to sample the phase space of jammed packings. This procedure samples each configuration proportionally to the volume of its basin of attraction.

**Hard-sphere fluid sampling.** We next take each equilibrated hard-disc fluid particle to diffuse at least a distance equal to the largest diameter in the system, as in ref. 13. To ensure statistical independence, we draw fluid configurations every $n_{\text{MC}}$ steps, where $n_{\text{MC}}$ is (pre-computed by averaging over multiple simulations) the total number of Markov chain Monte Carlo steps necessary for each individual particle to diffuse at least a distance equal to the largest diameter in the system.

**Soft shells and minimization.** We next take each equilibrated hard-disc fluid configuration and inflate the particles (instantaneously) with a WCA-like soft outer shell, to reach the target soft packing fraction $\phi_{\text{SS}} > \phi_{\text{HS}}$. Each hard sphere is inflated proportionally to its radius, so that the soft-sphere radius is given by

$$r_{\text{SS}} = \left( \frac{\phi_{\text{SS}}}{\phi_{\text{HS}}} \right)^{1/3} r_{\text{HS}}$$

where $d$ is the dimensionality of the box (2 in our case), and $r_{\text{HS}}$ and $r_{\text{SS}}$ are the soft- and hard-sphere radii respectively. Clearly, this procedure does not change the polydispersity of the sample. The radii are identical across volume fractions and system sizes, and the hard-disc fluid density is chosen so that the radius ratio of hard to soft discs is $(0.88/0.7)^{1/3} \approx 1.121$.

Next, particle inflation is followed by energy minimization using FIRE$^{15,12}$, to produce mechanically stable packings at the desired soft volume fractions $\phi$. This protocol has the advantage of generating packings sampled proportionally to the volume of their basin of attraction. In our simulations, we considered all mechanically stable packings, irrespective of the number of rattlers. To guarantee mechanical stability we required that the total number of contacts is sufficient for the bulk modulus to be strictly positive, $N_{\text{c,max}} = d(N_d - 1) + 1$ (ref. 33), where $N_{d}$ is the number of non-rattlers and $d$ is the dimensionality of the system.

Our implementation of FIRE enforces a maximum step size (set to be equal to $dN_{\text{c,max}}/2$) and forbids uphill steps by taking one step back every time the energy increases (and restarts the minimizer in the same fashion as the original FIRE implementation). We use a maximum time step $\Delta t_{\text{max}} = 1$, although the maximum step size is directly controlled in our implementation. All other parameters are set as in the original implementation$^{31}$.

**HS-WCA potential.** We define the WCA-like potential around a hard core as

$$v_{\text{HS-WCA}}(r) = \begin{cases} \infty & r \leq r_{\text{HS}}, \\ 4\epsilon \left( \frac{\sigma_{\text{HS}}}{r - 3\sigma_{\text{HS}}} \right)^{12} - \frac{\sigma_{\text{HS}}}{r - \sigma_{\text{HS}}} & r_{\text{HS}} < r < r_{\text{SS}}, \\ 0 & r \geq r_{\text{SS}}, \end{cases}$$

where $\sigma = (12\epsilon + \sigma_{\text{HS}}^3)^{1/3}$ guarantees that the potential function and its first derivative go to zero at $r_{\text{HS}}$. For computational convenience (avoids of square-root evaluations), the potential in equation (4) differs from the WCA form in that the inter-particle distance in the denominator of the WCA potential has been replaced with a difference of squares.

A power series expansion of equation (4) yields

$$v_{\text{HS-WCA}}(r) \approx 4\epsilon \left( \frac{12r_{\text{HS}}}{r_{\text{HS}}^2} \right)^{12} \left( r - r_{\text{HS}} \right)^2 + O \left( r - r_{\text{HS}} \right)^4$$

hence, in the limit of no overlap the pair potential is harmonic.

We numerically evaluate this potential, matching the gradient and linearly continuing the function $v_{\text{HS-WCA}}(r)$ for $r \leq r_{\text{HS}} + \epsilon$, where $\epsilon > 0$ is an arbitrary small constant, such that minimization is still meaningful if hard-core overlaps do occur. Our choice of potential is based on the fact that: the hard cores greatly reduce the amount of configurational space to explore, replacing expensive energy minimizations (to test whether the random walker has stepped outside the basin) with fast hard-core overlap rejections; and the hard cores exclude high-energy minima (jammed packings) that are not ‘hard-sphere-like’.

**Total accessible volume.** The basins of attraction of energy minima tile the ‘accessible’ phase space (schematically shown in Fig. 1b,c). This inaccessible part of the phase space arises due to hard-core constraints and the existence of fluid states

(see, for example, ref. 13). The total phase-space volume is equal to $V_{\text{box}}^\infty$. The inaccessible part of this volume arising from the hard-core constraints (shown as hatched areas in Fig. 1) is denoted by $V_{\text{in}}$, and the part corresponding to the coexisting unjammed fluid states is denoted by $V_{\text{SS}}$ (shown as blue regions with solid dots in Fig. 1b,c). $V_{\text{in}}$ is significant only for finite-size systems at or near unjamming. We denote the space tiled by the basins of mechanically stable jammed packings by $V_{\text{f}}$. We then have $V_{\text{f}} = V_{\text{box}}^\infty - V_{\text{in}} - V_{\text{SS}}$. In practice we compute $V_{\text{f}}$ using the following equation

$$\ln V_{\text{f}}(N, \phi_{\text{SS}}, \phi_{\text{HS}}) = \ln V_{\text{box}} - \ln N_{\text{HS}}(\phi_{\text{HS}}) + \ln \rho(\phi_{\text{HS}})$$

(6)

where $\rho(\phi_{\text{HS}})$ is the excess free energy, that is, the difference in free energy between the hard-sphere-fluid and the ideal gas, computed from the Santos–Yuste–Haró equation of state$^{31}$ as in ref. 13, and $\rho(\phi_{\text{HS}})$ is the probability of obtaining a jammed packing at soft volume fraction $\phi_{\text{HS}}$ with our protocol, shown in the inset of Fig. 3b.

**Counting by sampling.** We briefly review our approach to computing the number $\Omega$ of distinct jammed packings for a system of $N$ soft discs at volume fraction $\phi$. We prepare packings by the protocol described above, which generates jammed structures (energy minima) with probability $\rho(\phi_{\text{HS}})$ proportional to the volume of their basin of attraction $\nu_{\text{f}}$. We define the probability of sampling the $i$th packing as

$$p_i = \frac{\nu_i}{\nu_{\text{f}}}$$

(7)

where $\nu_i$ is the total accessible phase space, such that

$$\nu_{\text{f}} = \sum_i \nu_i$$

(8)

Details of the computation of $\nu_i$ are discussed in refs 13,16. To find $\Omega$, we make the simple observation

$$\sum_i \nu_i \Omega = \sum_i \frac{\nu_i \Omega}{\nu_{\text{f}}} \leq \sum_i \frac{\Omega}{\nu_{\text{f}}} \nu_i = \Omega$$

(9)

from which it follows immediately that

$$\Omega = \frac{\nu_{\text{f}}}{\nu_{\text{f}}}$$

(10)

The ‘Boltzmann-like’ entropy, suggested in a similar form by Edwards$, is then

$$S_\Omega = \ln \Omega = \ln N!$$

(11)

where the $\ln N!$ correction ensures that two systems in identical macrostates are in equilibrium under exchange of particles$^{17-19}$.

Note that $\nu(\phi)$ is the unbiased average basin volume (the mean of the unbiased distribution of volumes). We distinguish between the biased, $B(\phi; F)$ (as sampled by the packing protocol), and the unbiased, $U(\phi; F)$, basin log-volumes distributions ($F = -\ln \nu_{\text{f}}$). Since the configurations were sampled proportionally to the volume of their basin of attraction, we can compute the unbiased distribution as

$$U(\phi; F) = Q(\phi) B(\phi; F)$$

(12)

where $Q(\phi)$ is the normalization constant, such that

$$Q(\phi) = \int_{-\infty}^{\phi} dF B(\phi; F) e^{-F}$$

(13)

Since small basins are much more numerous than large ones, and grossly under-sampled, it is not sufficient to perform a weighted average of the sampled basin volumes. Instead, to overcome this problem, one can fit the biased measured basin log-volumes distribution $B(\phi; F)$ with an analytical (or at least numerically integrable) distribution, and perform the unbiased via equation (12) on the best fitting distribution. Different approaches to modelling this distribution give rise to different analysis methods, which all yield consistent results as shown in ref. 13. In this work we follow ref. 13 and fit $B(\phi; F)$ using both a (parametric) generalized Gaussian model$^{31}$ (see Supplementary Equation 21) and a (non-parametric) kernel density estimate with Gaussian kernels$^{31}$ and bandwidth selection performed by cross-validation$^{13,38}$, yielding consistent results in agreement with ref. 13.

Before performing the fit we remove outliers from the free-energy distribution in an unsupervised manner, as discussed in the ‘Data analysis’ section of the Methods.

No such additional steps are needed to compute the ‘Gibbs-like’ version of the configurational entropy, in fact

$$S_\Omega = \sum_i p_i \ln p_i - \ln N! = \sum_i [p_i (\ln v_i)] + \ln V_{\text{f}} - \ln N!$$

$$= (F)_B + \ln V_{\text{f}} - \ln N!$$

(14)

is simply the arithmetic average of the observed volumes: the sample mean of $F = -\ln \nu_{\text{f}}$ is already correctly weighted because our packing generation protocol generates packings with probability $p_i$. 

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Power-law between pressure and basin volume. A power-law relationship between the volume of the basin of attraction of a jammed packing and its pressure was first reported in ref. 13. In what follows we provide insight into this expression on the basis of this work’s findings. We observe that the structure of basins negative log-volumes, \( F = \ln v_{\text{basin}} \) and log-pressure, \( A = \ln P \), are approximately normally distributed (see Supplementary Figs 1 and 9). We therefore expect their joint probability to be well approximated by a bivariate Gaussian distribution \( B(\phi, F; A) \) following the distance-based outlier removal method introduced in ref. 43. This is applied in turn to each distribution, such that we choose to keep only those points for which at least \( R = 0.5 \) of the remaining data set is within \( D = 4 \varepsilon \) (matched with the much stricter \( R = 0.9988, D = 0.13 \sigma \) required to exclude any points further than \( |\mu - 3\sigma| \) for normally distributed data\(^4\)). On our data sets we find that this procedure removes typically none and at most 0.8% of all data points.

Mean and covariance estimates of \( B(\phi, F; A) \) are computed using a robust covariance estimator, namely the minimum covariance determinant (MCD) estimator\(^45\) with support fraction \( h_{\text{MCD}} = 0.99 \). The MCD estimator defines \( \mu_{\text{MCD}} \), the mean of the \( h \) observations for which the determinant of the covariance matrix is minimal, and \( \sigma_{\text{MCD}} \), the corresponding covariance matrix\(^46\). We use these robust estimates of the location and of the covariance matrix (computed over 1,000 bootstrap samples\(^{47} \)) to fit our observations by linear minimum mean square error\(^2\) (see Fig. 2).

Before fitting \( B(\phi) \) (required to compute \( \mu \)), we perform an additional step of outlier detection based on an elliptic (Gaussian) envelope criterion constructed using the MCD estimator. We assume a support fraction \( h_{\text{MCD}} = 0.99 \) and a contamination equal to 10% (ref. 37). We compute \( S_8 \) and \( S_9 \) from the resulting data sets. The procedure is strictly unsupervised and allows us to achieve robust fits despite the small sample sizes. We fit \( B(\phi) \) using both a (parametric) generalized Gaussian model\(^47\) and a (non-parametric) kernel density estimate with Gaussian kernels\(^48,49\) and bandwidth selection performed by cross-validation\(^5\). 

Data availability. The data that support the plots within this paper and other findings of this study are available from the University of Cambridge data repository doi: http://dx.doi.org/10.17863/CAM.9985.

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