PHYSICAL REVIEW E 95, 052905 (2017)

Affine and topogical structural entropies in granular statistical mechanics: Explicit calculations and equation of state

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(Received 23 January 2017; published 31 May 2017)

We identify two orthogonal sources of structural entropy in rattler-free granular systems: affine, involving structural changes that only deform the contact network, and topological, corresponding to different topologies of the contact network. We show that a recently developed connectivity-based granular statistical mechanics separates the two naturally by identifying the structural degrees of freedom with spanning trees on the graph of the contact network. We extend the connectivity-based formalism to include constraints on, and correlations between, degrees of freedom as interactions between branches of the spanning tree. We then use the statistical mechanics formalism to calculate the partition function generally and the different entropies in the high-angoricity limit. We also calculate the degeneracy of the affine entropy and a number of expectation values. From the latter, we derive an equipartition principle and an equation of state relating the macroscopic volume and boundary stress to the analog of the temperature, the contactivity.

DOI: 10.1103/PhysRevE.95.052905

I. INTRODUCTION

Granular matter is one of the most significant forms of matter in nature, both on Earth and celestially. It is also relevant to human society in many ways, be it in the context of products and technological applications or through our interactions with the natural environment around us. Yet, a reliable fundamental understanding of this form of matter is yet to emerge, a situation that is limiting the development of predictive and effective modeling. Consequently, this area has been the focus of intensive research in recent years.

One of the main modeling tools in the theorist's arsenal is statistical mechanics. This powerful method, devised originally to deal with thermodynamic systems subject to thermal fluctuations, is the ultimate coarse-graining technique. Statistical studies of granular assemblies date back to the 1920s [1], but the introduction of granular statistical mechanics (GSM) in 1989 [2,3] led to a significant increase in research in this direction within the physics community. A number of major advances include the calculation of the volume function [4], the introduction of the stress ensemble [5,6], the finding that the two ensembles are interdependent [7], and the measurement of either ensemble's equilibration [8]. The potential advantage of statistical mechanics is the ability to derive with it equations of state and constitutive relations, a long-sought goal in the field. Yet, such relations have been slow to emerge for a variety of reasons [9-11].

GSM is based on entropy, namely the number of structural and stress configurations that static assemblies of macroscopic particles can have [12–14]. Recently, the entropy of packs of up to N = 128 soft particles was measured numerically and found to be extensive after the subtraction of $\ln N!$ [15–17].

In general, GSM consists of two subensembles, one of all the structural configurations and the other of all stress microstates. Originally, the structural microstates were proposed to occur with probability that depends on their volume [2,3], but it was shown recently that this formulation is flawed [18], which may have also been responsible for the little use of the GSM in the community. Another formulation, based on a connectivity function, was then proposed, but it has not been tested yet. The connectivity-based partition function is

$$Z = \int \prod_{n=1}^{N_c - 1} d\vec{r}_n \prod_{m=1}^M d\vec{g}_m \Theta(\{\vec{r}\}) G(\{\vec{r}\}) \\ \times e^{-\frac{C(\vec{r})}{\tau} - \gamma: \mathcal{F}(\{\vec{r}\}, \{\vec{g}\})},$$
(1)

where the vectors $\{\vec{r}\}$ connect contacts around particles and $\{\vec{g}\}$ are the compressive forces acting on the boundary particles [see Fig. 1 for an example in two dimensions (2D)]. $\tau =$ $\partial \langle \mathcal{C} \rangle / \partial S$, with S the entropy, is the contactivity, a measure of the connectivity fluctuations that is an analog of the temperature. \mathcal{F} is the force moment tensor, formed by the outer product of the intergranular forces and their position vectors, summed over all contacts. This function couples the structure and the stress ensembles [5–7]. $\gamma_{\alpha\beta} = 1/X_{\alpha\beta}$ is an (inverse) angoricity tensor [5] and Θ includes the constraints on the systems forming the ensemble, e.g., that they are in mechanical equilibrium, generated by the same process and all have the same mean coordination number \bar{z} . G is called the measure, which is independent of the exponential Boltzmann factor and represents the probability to sample a specific configuration of vectors \vec{r} . The measure may depend on the preparation and sampling protocols, and need not be uniform, as recent results suggest [15–17]. Its form is not known for our systems and, for simplicity, we take it to be uniform. Nevertheless, the following analysis can be carried out for any form of G: The connectivity, $C = \sum_{q} \vec{r}^{q} \cdot \vec{r}^{q}$, is a sum over all N_{r} intercontact vectors. Of these vectors, only $N_c - 1$ are independent, where N_c is the number of contacts. In principle, the structural degrees of freedom (DFs) should also include the parameters specifying the particle shapes. These are ignored here for brevity but could be included without loss of generality [11].

In this paper, we demonstrate the use of the full structurestress partition function, whose structural part is based on the recent connectivity function. The paper is constructed as follows: In Sec. II, we rewrite the partition function of Eq. (1) in a convenient form. In Sec. III, we clarify the distinction between two different types of entropies in granular materials: affine and topological. In Sec. IV, we calculate expectation values and the affine entropy. We also derive an equipartition principle and an equation of state. In Sec. V, we show that the connectivity partition function readily accommodates structural constraints and correlations between the vectors \vec{r} . In Sec. VI, we outline the calculation of the topological entropy. We conclude in Sec. VII.

II. THE PARTITION FUNCTION

The vectors \vec{r} are the structural DFs. In 2D, they run clockwise around each particle (see Fig. 1) and there are $N\bar{z}$ of them. A similar parametrization exists in three dimensions (3D) [13,19]. For clarity, we focus here on 2D systems experiencing no body forces. We denote by M the number of boundary particles to which external compressive forces are applied and by $(\alpha - 1)M$ [$\alpha = O(1)$] the total number of boundary particles. It follows that αM is the number of the outermost boundary vectors \vec{r} in the system.

In 2D, the vectors \vec{r} form loops around particles and around cells and, therefore, only $N_{\rho} = N_c - 1$ of them are independent. The set of independent vectors \vec{r} , of which there are many possibilities, forms an undirected spanning tree on the contact network [20]. As we will see below, it is convenient to constrain our choice of spanning tree to include $\alpha M - 1$ of the outermost boundary vectors and denote those by \vec{r}^b . For brevity, we define \vec{R} as the $(d \times N_r \log)$ vector, containing the components of all the vectors \vec{r} . The entries of \vec{R} are ordered as follows: First come the $\alpha M - 1$ independent boundary vectors \vec{r}^b (*d* entries each), then $N_c - \alpha M$ independent vectors \vec{r} in the bulk, and then the remaining (dependent) vectors \vec{r} . We can then write

$$\vec{R} = A\vec{\rho},\tag{2}$$

where $\vec{\rho}$ is a $d \times N_{\rho}$ long vector, containing only the components of the independent vectors \vec{r} . The top part of A is clearly a unit matrix, whereas each line of the bottom part describes the route along the spanning tree to get from the tail to the head of a certain dependent \vec{r} . It follows that all the entries of A are predominantly 0, with fewer ± 1 entries. In terms of $\vec{\rho}$ we have

$$C/\tau = \vec{\rho} A^T A \vec{\rho} / \tau = \frac{1}{2} \vec{\rho} B \vec{\rho} \qquad \left(B \equiv \frac{2}{\tau} A^T A \right). \tag{3}$$

The stress ensemble is controlled by the force moment tensor, which can be written either as a sum over the outer product of the contact force and position at every contact point, or in terms of the loop forces, \vec{f}^i , defined in Ref. [21], namely, $\mathcal{F}_{\alpha\beta} = \sum f_{\alpha}^i r_{\beta}^i$. While the sum is over all the cells, the contribution of internal cells vanishes. This is because \vec{f}^i is constant for every cell, and the sum over the vectors \vec{r} that enclose the cell vanishes. It follows that the only contribution to \mathcal{F} comes from the boundary vectors, $\sum_{b=1}^{\alpha M} f_{\beta}^b r_{\beta}^b$. These vectors are all, but one, independent and constitute the first entries of $\vec{\rho}$. The loop forces \vec{f}^b are accumulation of the boundary forces \vec{g}^b (see Fig. 1). Recalling that the loop force, associated with the single dependent boundary vector, to zero. We can thus manipulate the force moment term in the partition function to the form

$$\gamma : \mathcal{F} = \vec{g}H^T \gamma E \vec{\rho} = \vec{g}Q^T \vec{\rho} \quad (Q \equiv E^T \gamma^T H).$$
(4)

(5)

	(0)	0	0	0	0		0	0)	
	1	0	0	0	0		0	0	
	1	1	0	0	0		0	0	
H =	1	1	1	0	0		0	0	;
	:	÷	÷	÷	÷	۰.	÷	÷	
	$\backslash 1$	1	1	1	1		1	0/	

	boundary		boundary		boundary			boundary			bulk	
	cell 1		cell 2		cell 3			cell $M-1$			vectors	
$E = \begin{pmatrix} 0 \\ 1 \\ 0 \\ 0 \\ 0 \\ 0 \end{pmatrix}$	0 1 0 0 :	0 1 0 0 : 0	0 1 0 0 :	$ \begin{array}{c} -1 \\ 0 \\ 1 \\ 0 \\ \vdots \\ 0 \end{array} $	$ \begin{array}{c} -1 \\ 0 \\ 1 \\ 0 \\ \vdots \\ 0 \end{array} $	$ \begin{array}{c} -1 \\ 0 \\ 0 \\ 1 \\ \vdots \\ 0 \end{array} $	$ \begin{array}{c} -1 \\ 0 \\ 0 \\ 1 \\ \vdots \\ 0 \end{array} $	$ \begin{array}{c} -1 \\ 0 \\ 0 \\ 1 \\ \vdots \\ 0 \end{array} $		$\begin{vmatrix} -1 \\ 0 \\ 0 \\ \vdots \\ 1 \end{vmatrix}$	$ \begin{array}{c} -1 \\ 0 \\ 0 \\ \vdots \\ 1 \end{array} $	0



FIG. 1. A 2D polydisperse granular pack with $(\alpha - 1)M = 18$ boundary particles (slightly shaded), of which M = 12 touch the walls (and thus $\alpha = 2.5$). The vectors \vec{r} (solid and dashed, thick and thin) connect a particle's nearest contacts, circulating clockwise. Their components constitute the vector \vec{R} . The solid vectors (both thick and thin) form a nondirectional spanning tree: They form no loops, representing the independent DFs, and they reach every contact point. Their components constitute the vector $\vec{\rho}$. Each dashed vector is a linear combination of solid vectors, namely one can always get from its tail to its head by walking along solid vectors. There are $\alpha M = 30$ boundary (thick) vectors \vec{r} , and our choice of spanning tree includes all of them but one. Overall the pack has $N_c = 56$ contacts and thus $N_{\rho} = 55$ independent (solid) vectors \vec{r} . Also illustrated are three boundary forces, \vec{g}^b , and three boundary loop forces, \vec{f}^b . As explained in the text, we set $\vec{f}^0 = 0$, and the rest are accumulation of the boundary forces, going counterclockwise around the pack.

The matrices *E* and *H*, that correspond to the system in Fig. 1, are defined in Eq. (5) below. *H* is the accumulation matrix, transforming from the boundary forces, \vec{g} , into the loop forces, $\vec{f} = H\vec{g}$, where we set $\vec{f}^0 = 0$. *E* transforms from the independent vectors, $\vec{\rho}$, into the boundary vectors, \vec{r}^b , and sums the boundary vectors \vec{r}^b of each boundary cell. These are always several consecutive vectors from the *M* – 1 first entries of $\vec{\rho}$. Since the first cell includes one dependent boundary vector, the sum of this cell's vectors is the (negative) sum of all the other boundary vectors \vec{r}^b . Noting that the matrices *A*, *B*, *E*, and *H* do not involve the Cartesian coordinates of the matrix γ , the latter could be placed as shown in (1). This position is arbitrary with respect to *H* or E^T in the definition of *Q*.

III. AFFINE AND TOPOLOGICAL STRUCTURAL ENTROPIES

Before we continue, it is essential to identify and classify three different sources contributing to the entropy of granular systems, (i) S_r is the *rattler entropy*, which consists of all the positional variations of the rattlers inside voids. The rattlers are particles that, in the absence of external and body forces, are not part of the force-carrying structure. This contribution is essentially the volume that these rattlers can occupy within any given structure and as such it is straightforward to calculate. Nevertheless, it has little bearing on most of the mechanisms governing the physics of static granular assemblies, which involve force transmission between particles. Therefore, this contribution is ignored in the following discussion.

(ii) S_a is the *affine entropy*. Regarding the contact network as vertices of a graph, whose edges are the vectors \vec{r} , S_a consists of all the possible affine, i.e., connectivity-preserving, distortions of this graph, which the given collection of N force-carrying particles can make. We shall include the entropy coming from the stress DFs in S_a as well.

(iii) S_t is the *topological entropy*, which consists of all the possible different topologies, or connectivity networks, that the same collection of particles can make.

Equation (6) is based on the assumption that the three entropies are additive, i.e., that the partition function is a product of three independent parts, $Z = Z_r Z_a Z_t$. This mean-field-like approximation is justified in Sec. VI. Note that the distinction between S_a and S_t is useful mainly when the mean coordination number is kept fixed across the systems in the ensemble, which is implicitly assumed here. Letting it fluctuate gives rise to different topologies, obviating the affine entropy. As we will show below, the distinction between these two types of entropy is crucial because their scaling with N is markedly different.

IV. AFFINE ENTROPY: CALCULATION, EXPECTATION VALUES, AND EQUATION OF STATE

All the connectivity, or topological, information is contained in the matrix A in the sense that the connectivity of a specific configuration corresponds to a specific set of entries A_{ij} . Thus, S_a is the result of the different changes in the vectors \vec{r} , under the constraint that the matrix A, and hence B, are fixed. To calculate the affine entropy, we use Eqs. (3) and (4) to express the partition function of Eq. (1) in terms of $\vec{\rho}$ and \vec{g} :

$$Z_{a} = \int e^{-\frac{1}{2}\vec{\rho}B\vec{\rho} - \vec{g}Q^{T}\vec{\rho}} (d\vec{\rho}) (d\vec{g}).$$
(7)

In the following, we first calculate Z_a while ignoring the constraints and taking $\Theta = 1$. We will then point out the problems that this simplification leads to and adjust the calculation to model in the constraints. The integral in (7) is Gaussian and its calculation straightforward. Changing variables, $\vec{\rho} \equiv \vec{\rho} + B^{-1}Q\vec{g}$, we have

$$-\frac{1}{2}\vec{\rho}B\vec{\rho} - \vec{g}Q^{T}\vec{\rho} = -\frac{1}{2}\vec{\tilde{\rho}}B\vec{\tilde{\rho}} + \frac{1}{2}\vec{g}P\vec{g},$$
(8)

with $P \equiv Q^T B^{-1} Q$. The partition function then reduces to two decoupled integrals,

$$Z_{a} = \int e^{-\frac{1}{2}\vec{\rho}B\vec{\rho}}(d\vec{\rho}) \int e^{+\frac{1}{2}\vec{g}P\vec{g}}(d\vec{g}),$$
(9)

which can be calculated readily because *B* and *P* are symmetric and hence diagonalizable by orthogonal matrices. *B* is positive definite, since for any choice of $\vec{\rho} \neq 0$ we have

 $\vec{\rho} B \vec{\rho} = \vec{R} \cdot \vec{R} > 0$, which means that all its eigenvalues are positive. The matrix *P*, however, is singular—it has *d* zero eigenvalues, leading to an integrand of 1. The other eigenvalues are positive, leading to an *increasing exponential* due to the positive sign in Eq. (9). This, however, does not lead to a diverging integral because the boundary forces, \vec{g} , are finite. The integrals in (9) are straightforward to calculate,

$$Z_a = \sqrt{\frac{(2\pi)^{dN_p} 2^{3d(M-1)}}{|B|^d |P|_+}} (2g)^d e^{\frac{g^2}{2} \operatorname{Tr}(P)} \prod_{p_i > 0} D(a_i), \quad (10)$$

where |B| is the determinant of B, p_i are the eigenvalues of P, $|P|_+$ is its pseudodeterminant (i.e., the product of all nonzero p_i 's), g is the maximal boundary force, $a_i \equiv g\sqrt{p_i/2}$, and D is the Dawson function:

$$D(\alpha) \equiv e^{-\alpha^2} \int_0^\alpha e^{x^2} dx.$$
 (11)

We can now use Eq. (10) to calculate the affine entropy and several expectation values, for a given contact network. The detailed calculations are shown in Appendix A and here we present the main results.

(1) The mean connectivity can be calculated using $\langle C \rangle_a = \tau^2 (\partial \ln Z_a / \partial \tau)$, while keeping A constant,

$$\langle \mathcal{C} \rangle_a = \frac{\tau}{2} \left[d(N_\rho - M + 1) + \sum_{a_i > 0} \frac{a_i}{D(a_i)} \right].$$
(12)

In the high angoricity limit, $a_i/D(a_i) = 1$, and since there are d(M - 1) of them, Eq. (12) becomes

$$\langle \mathcal{C} \rangle_a = N_\rho d\tau / 2 \,. \tag{13}$$

This is an equipartition principle [18]: The mean connectivity is shared among the dN_{ρ} structural DFs, with each getting on average $\tau/2$, analogously to the mean energy of $k_BT/2$ per DF in thermal systems. Equation (12) deviates from this by $O(M \sim \sqrt{N} \ll N)$, since the sum on the right is over d(M - 1) elements of O(1).

In the low angoricity limit, $D(a_i) \approx 1/2a_i$, and the sum on the right-hand side of (12) becomes $g^2 Tr(P)$. This is a sum over O(M) finite terms and it scales as $\tau \operatorname{Tr}(\gamma^2)$. Thus, it is also negligible relative to the first term as long as $g^2 \tau \operatorname{Tr}(\gamma^2) < O(M)$, and equipartition holds in this regime too up to terms of $O(\sqrt{N})$.

(2) The mean squared norm of the force vector is

$$\langle \vec{g} \cdot \vec{g} \rangle_a = -\operatorname{Tr}(\Pi) + \sum_{a_i > 0} \frac{g^2}{2a_i D(a_i)},\tag{14}$$

where Π is the pseudoinverse matrix of P, $P\Pi P = P$. Note that $P\Pi \neq I$, as P is singular. The statistics of the boundary forces should be hardly dependent on the precise internal topology and, hence, $\langle \vec{g} \cdot \vec{g} \rangle_a = \langle \vec{g} \cdot \vec{g} \rangle$. The first term on the right-hand side scales as $1/[\tau \operatorname{Tr}(\gamma^2)]$. The dependence of the second term on τ and γ is more complex as it is via the nonlinear Dawson function. However, this dependence can be obtained in two limits. In the high angoricity limit $D(a_i) \approx$ a_i and the dependence is the same, $1/[\tau \operatorname{Tr}(\gamma^2)]$. In the low angoricity limit, $D(a_i) \approx 1/2a_i$, the second term approaches a constant and its dependence on τ and γ disappears.

(3) Taking the system to be a square of $\sqrt{Na} \times \sqrt{Na}$, where *a* is the typical size of a particle, and assuming isotropic boundary stresses, the total force normal to one side is $M\langle g_x \rangle_a/4$, with g_x the component of a boundary force normal to the wall. An explicit calculation of the expectation value $\langle g_x \rangle_a$ (see Appendix A) gives $\langle g_x \rangle_a = |\mathcal{G}|/\sqrt{M}$, with \mathcal{G} an *M*-long vector, whose components are $\mathcal{G}_i = g(1 - e^{-a_i^2})/[2a_i D(a_i)]$. Summing over all the normal components of the external forces along the side and dividing by the length, we obtain the expectation value of the normal stress

$$\langle \sigma_{nn} \rangle_a = \frac{M}{4\sqrt{Na}} \frac{|\mathcal{G}|}{\sqrt{M}} \approx \frac{|\mathcal{G}|}{4a\sqrt[4]{N}}.$$
 (15)

The components of \mathcal{G} range between 0.5 and 1, as we show explicitly in Appendix C, leading to $|\mathcal{G}| = O(\sqrt{M} \approx \sqrt[4]{N})$. Therefore, $\langle \sigma_{nn} \rangle_a$ is independent of system size, as expected.

(4) The affine entropy, given by $S_a = \langle C \rangle_a / \tau + \ln Z_a$, is

$$S_{a} = -\frac{d}{2} \ln |B| - \frac{1}{2} \ln |P|_{+} + \frac{g^{2}}{2} \operatorname{Tr}(P) + \sum_{a_{i}>0} \left[\ln D(a_{i}) + \frac{a_{i}}{2D(a_{i})} \right] + \frac{d}{2} \left[N_{\rho} \ln(2\pi e) + 2\ln(2g) + (M-1) \ln\left(\frac{8}{e}\right) \right].$$
(16)

There is more to this result than being satisfyingly exact and testable: It also provides the following significant observation. The terms involving |B| and N_{ρ} are of order N, thus dominating over all the other terms, which are of order $M \sim \sqrt{N}$. Since these two terms originate only in the structure, they are independent of the external boundary forces. This means that, for sufficiently large systems, *the boundary forces contribute negligibly to the affine entropy regardless of the angoricity value*. Consequently, Eq. (16) substantiates the generality of previous results obtained in the high angoricity limit. Keeping only these two terms, we obtain

$$S_a \approx \frac{d}{2} [N_\rho \ln(2\pi e) - \ln |B|].$$
 (17)

This result was also derived and calculated numerically in Ref. [18]. In particular, it was shown there to scale linearly with N. This makes the affine entropy conveniently extensive. It also increases with τ , the measure of structural fluctuations, as expected. In Appendix B we discuss in more detail different approximations for S_a .

(5) To calculate the mean volume, note that the volume is a quadratic function of all the independent vectors \vec{r} , $V \equiv \vec{\rho} W \vec{\rho}$. Rewriting it in terms of the transformed variables of Eq. (8), we have

$$V = \vec{\rho} W \vec{\rho} = \vec{\tilde{\rho}} W \vec{\tilde{\rho}} - 2\vec{g} Q^T B^{-1} W \vec{\tilde{\rho}} + g U g, \qquad (18)$$

with $U \equiv Q^T B^{-1} W B^{-1} Q$. The cross term vanishes on integration over the symmetric distribution of $\vec{\rho}$, giving

$$\langle V \rangle_{a} = \frac{1}{Z_{a}} \int (\vec{\rho} W \vec{\rho} + g U g) e^{-\frac{1}{2}\vec{\rho} B \vec{\rho} + \frac{1}{2} \vec{g} P \vec{g}} (d \vec{\rho}) (d \vec{g})$$

= $d \operatorname{Tr}(W B^{-1}) - \operatorname{Tr}(U \Pi) + \sum_{a_{i} > 0} \frac{g^{2} U_{ii}'}{2a_{i} D(a_{i})},$ (19)

with U' being the matrix U, rotated to the basis where P is diagonal (see Appendix A for details). The first two terms in Eq. (19) are linear in τ and are independent of γ , but the third term depends on both τ and γ . As in Eq. (14), this dependence is not simple. However, in the high angoricity limit it becomes also linear in τ , and at low angoricity it scales as $\tau^2 \operatorname{Tr}(\gamma^2)$. The expectation value $\langle V \rangle_a$ suffers from a problem, which we discuss in detail and then resolve in the next section.

(6) To obtain the affine contribution to the equation of state, we combine Eqs. (15) and (19) to obtain

$$\langle \sigma_{nn} \rangle_a \langle V \rangle_a = F(\tau, X_{\alpha\beta}, g).$$
 (20)

The form of this expression resembles that of the ideal gas equation of state, $PV = Nk_BT$: The left-hand side involves two macroscopically measurable quantities, while the right-hand side is a function of the temperature-like variables alone, the angoricity and contactivity.

The above expectation values and equation of state, (12), (14), (15), (16), (19), and (20), can be simplified by approximating the Dawson function for low and high angoricities, as we show in detail in Appendix B. In particular, in the high angoricity limit, we obtain the following explicit equation of state:

$$\langle \sigma_{nn} \rangle_a \langle V \rangle_a \approx \frac{g}{8} M \bigg[d \operatorname{Tr}(WB^{-1}) + \frac{g^2}{3} \operatorname{Tr}(U\Pi P) \bigg].$$
 (21)

In this expression, the first term on the right-hand side scales as τ and the second term scales as $\tau^2 \operatorname{Tr}(\gamma^2)$. In this limit, the second term, which originates in the forces entropy, is negligibly small compared to the first, in agreement with our observation, Eq. (16).

V. EXTENDING THE FORMALISM: INCLUDING CONSTRAINTS AND VECTOR-VECTOR INTERACTIONS

A close scrutiny of the result for $\langle V \rangle_a$, Eq. (19), reveals a problem: The expected volume vanishes. To see this, we introduce a projection operator onto the Cartesian coordinates, \mathcal{J} . Using this operator on B, which does not depend on the Cartesian coordinates, gives $\mathcal{J}(B) = 1$, with 1 the 2D identity matrix; $\mathcal{J}(W) = \epsilon$, with ϵ the 2D Levi-Civita symbol, since volume is a sum of cross products; $\mathcal{J}(P) = \gamma \gamma^T$; and $\mathcal{J}(U) =$ $\gamma \epsilon \gamma^T$. Using these, the first term on the right-hand side of Eq. (19) vanishes, $\text{Tr}(WB^{-1}) \propto \text{Tr}(\mathcal{J}(WB^{-1})) = \text{Tr}(\epsilon) = 0$. The second term also vanishes because $\text{Tr}(U\Pi) \sim \text{Tr}(\epsilon) = 0$. Similar considerations give that $U'_{ii} = 0$, which means that the third term also vanishes and we obtain that overall $\langle V \rangle_a = 0$.

This, of course, cannot be correct. The problem can be traced to the omission of the function Θ from the partition function, which allowed the independent vectors, \vec{r} , to be unconstrained in the integral (19). In particular, for every occurrence of \vec{r} there is an occurrence of $-\vec{r}$. This means

that, for every structural configuration we consider, there is a configuration with a volume of the opposite sign, which cancels its contribution to the integral.

Indeed, the integration over the independent vectors \vec{r} cannot be unconstrained since they must not cross one another (see, e.g., Fig. 1). To accommodate this condition in two dimensions, we constrain the vectors to rotate always clockwise around particles and anticlockwise around cells. This can be implemented by ensuring that the cross-product of successive vectors must be negative in circulating around particles and positive in circulating around cells. Thus, defining the trend function,

$$T(\vec{r}^i, \vec{r}^j) \equiv \vec{r}^i \times \vec{r}^j, \tag{22}$$

we introduce the following constraints into Z_a :

$$\Theta \equiv \prod_{g}^{\text{grains}} \prod_{i=1}^{z_g} \mathcal{H}[-T(\vec{r}^{(g,i)}, \vec{r}^{(g,i+1)})]$$

$$\cdot \prod_{c}^{\text{cells}} \prod_{i=1}^{z_c} \mathcal{H}[T(\vec{r}^{(c,i)}, \vec{r}^{(c,i+1)})], \qquad (23)$$

with \mathcal{H} being the Heaviside step function. To satisfy these constraints, we augment the connectivity function by introducing a set of Lagrange multipliers, $\lambda^{(g/c,i)}$:

$$C = C_0 - \sum_{g,i} \lambda^{(g,i)} T(\vec{r}^{(g,i)}, \vec{r}^{(g,i+1)}) + \sum_{c,i} \lambda^{(c,i)} T(\vec{r}^{(c,i)}, \vec{r}^{(c,i+1)}).$$
(24)

An advantage of this formulation is that it does not complicate the calculation. Our added constraints involve only quadratic terms in $\vec{\rho}$ and, therefore, they only modify the numerical values of the entries of the matrix *B* in the partition function. It follows that our results, Eqs. (12)–(20) are still valid, but with a modified matrix $B \rightarrow B_T$. Using our projection operator, $\mathcal{J}(T) = \epsilon$ and, hence, $\mathcal{J}(B_T) \neq 1$, resolving the problem of the vanishing mean volume in Eq. (19).

We now note that a similar tactic can be employed to take into account correlations between vectors \vec{r} , which are inherently present in real granular packs. For example, the angle, θ_{ij} , between two successive vectors, \vec{r}^i and \vec{r}^j , assumes very specific values, given the order of the particle or cell that they share. To demonstrate this, we analyze 396 computergenerated systems of 64 soft particles [16]. The particle radii are normally distributed, with a mean of 1 and a standard deviation of 0.1 in arbitrary units. The systems were generated at a constant volume fraction, $\phi = 0.84$, and their mean coordination numbers range between 4 and 4.4 (discarding rattlers). Figure 2 shows a histogram of θ_{ij} , according to cell and particle order. Given the order of the particle or cell, θ_{ij} is narrowly distributed around a certain value, $\mu_{ij}^{(\theta)}$, with a certain standard deviation, $\sigma_{ij}^{(\theta)}$. Using a combination of dot products and cross-products, we can construct potential wells around these preferred values. Concretely, by adding the following



FIG. 2. The histograms of θ_{ij} , for 396 systems of 64 particles each, in two cases: (a) when \vec{r}^i and \vec{r}^j share a particle and (b) when they share a cell. The colors red, yellow, green, and blue correspond to a particle and cell of order 3, 4, 5, and 6, respectively. The angle in cells of order 3, for example, is narrowly distributed around $\pi/3$, corresponding to an equilateral triangle. The deviation from $\pi/3$ is due to the distribution of particle sizes.

term to the connectivity,

$$\mathcal{C}_{ij}^{(\theta)} = a(\vec{r}^i \cdot \vec{r}^j) + b(\vec{r}^i \times \vec{r}^j)$$
$$= |\vec{r}^i| |\vec{r}^j| (a\cos\theta_{ij} + b\sin\theta_{ij}), \qquad (25)$$

we create a potential well around $\mu_{ij}^{(\theta)} = \arctan(b/a)$. The magnitudes of *a* and *b* determine the depth of the well and thus determine $\sigma_{ij}^{(\theta)}$. Such a term can be added for every two successive vectors, with the appropriate parameters *a* and *b*. These additions, like the augmented connectivity of Eq. (24), are all quadratic in $\vec{\rho}$, and thus do not complicate the calculation.

The addition of such terms introduces explicitly interactions between the DFs, a concept that has been absent from the original formulation of GSM.

Another example of an interaction between DFs is the inherent correlation between the lengths of successive vectors \vec{r} around a particle. Figures 3(a) and 3(b) show the histogram of $|\vec{r}|$, sampled in two different ways: one by the order of its cell [Fig. 3(a)] and the other by the order of its particle [Fig. 3(b)]. We see that $|\vec{r}|$ increases with cell order and decreases with particle order. This has a geometrical origin-the more vectors surround a particle, the shorter they must be, but the more vectors constitute a cell, the longer they can be (see Fig. 1). Let us focus on order 3 cells, the tall peak in Fig. 3(a). Figures 3(c)and 3(d) show, respectively, 2D histograms (shown as heat maps) of the expected correlation-free and actual occurrence frequencies of the lengths of successive vectors, \vec{r}^i and \vec{r}^j , around order 3 cells. The differences between the two indicate a positive correlation between the lengths, which is the result of a correlation in the lengths of the three vectors upon increase of the size of any of the three particles around the cell.

The same can be done for particles of any number of contacts or cells of any order. In Figs. 3(e) and 3(f), we show, respectively, the expected correlation-free and actual distributions for order 4 cells. Here too the two differ significantly, but with a negative correlation that arises from a geometrical origin: The lengths of two opposite edges in order 4 cells are sensitive to the distance between the two respective particles (see Fig. 1): the longer one pair of opposite edges, the shorter the other.



FIG. 3. [(a), (b)] The histogram of $|\vec{r}|$, for 396 systems of 64 particles each, in units of the average particle radius. Each vector \vec{r} belongs to one particle and one cell, and the histogram is divided according to the order of the cell (a) and the particle (b), with orders 3, 4, 5, and 6 represented by red, yellow, green and blue, respectively. [(c), (e)] The expected 2D histogram of successive vectors, \vec{r}^i and \vec{r}^j , around order 3 and 4 cells, respectively, according to panel (a), and not taking correlations into account. [(d), (f)] The actual 2D histogram corresponding to panels (c) and (e). The heat maps on the right differ from those on the left due to positive correlations for order 3 cells and negative correlations for order 4 cells.

To take account of such correlations, we can include in the connectivity function another interaction-like term between two successive vectors \vec{r}^i and \vec{r}^j :

$$\mathcal{C}_{ii}^{|r|} = (|\vec{r}^i|^2 - |\vec{r}^j|^2)^2, \tag{26}$$

which is quartic in $\vec{\rho}$. The calculation of the partition function with quartic terms is more complex and is left for a future study.

VI. THE TOPOLOGICAL ENTROPY

The topological entropy arises from all the possible changes in the contact network, i.e., the topology. Each such change, which corresponds to making or breaking a contact, effects a change in the matrix A. The inverse, however, is not generally true—many different matrices A may correspond to the same topology because the choice of the spanning tree is not unique. Thus, to obtain the number of possible topologies, we need to divide the total number of possible matrices A, Ω_t , by the number of spanning trees describing a typical configuration, N_{ST} . We call the latter multiplicity of a microstate.

The statistics of A can be described by a multivariate probability density function of its components, $P(\{A_{ij}\})$. The partition function is then

$$Z = \sum_{A_{ij}} P(\{A_{ij}\}) Z_a(A),$$
(27)

where the sum is over all possible values of A_{ij} .

To estimate $P(\{A_{ij}\})$, we first recall that the top part of A, corresponding to the N_{ρ} independent vectors, is always a unit matrix. The rest of the matrix consists of entries of 0 and ± 1 , whose ij positions can change from topology to topology. A straightforward naive approximation is to assume that each A_{ij} is independent and can take the values 0, 1, and -1 with respective probabilities P_0 , P_1 , and $P_{-1} = 1 - P_0 - P_1$. These probabilities are constrained by the requirement that $P_1 + P_{-1} = \langle l \rangle / N_{\rho}$, where $\langle l \rangle$ is the average number of steps along the spanning tree needed to describe a dependent vector \vec{r} in terms of the independent ones. The average $\langle l \rangle$ is carried out over all possible topologies (possible matrices A) generated by the same packing procedure of a specific collection of N particles, resulting in the same mean coordination number \bar{z} .

However, this estimate can be improved. Since each line of *A* represents a route along the spanning tree, a more realistic estimate is

$$P(\{A_{ij}\}) = \prod_{i} P(n_0^{(i)}, n_1^{(i)}, n_{-1}^{(i)}),$$

with $n_0^{(i)}$, $n_1^{(i)}$, and $n_{-1}^{(i)}$, respectively, being the numbers of 0's, 1's, and -1's in the *i*th row of *A*. Different topologies correspond to different bottom parts of *A*, consisting of $N_r - N_\rho \sim 2N_c - N_c = N_c$ rows and $N_\rho \sim N_c$ columns. Typically, each of those N_c rows has $n \equiv \langle l \rangle / 2$, of 1's and -1's. Therefore, with $n \ll N_c$, the total number of possible different *A* matrices is

$$\Omega_t \approx \left[\binom{N_c}{n} \binom{N_c}{n} \right]^{N_c} \approx \left(\frac{N_c^n}{n!} \right)^{2N_c}, \qquad (28)$$

and we have

$$\ln \Omega_t = 2N_c [n \ln N_c - \ln n!] + O(n).$$
(29)

We now need to estimate the multiplicity of spanning trees per topology, N_{ST} . This can be done using a result from graph theory: For a wide class of graphs [22],

$$\lim_{N_V \to \infty} N_{\rm ST}^{(G)} = e^{\zeta_G N_V},\tag{30}$$

where ζ_G is a constant, whose value depends on the particular topology of the graph G, and N_V is the number of the graph's vertices, which is N_c in our case. A straightforward upper bound on N_{ST} can be established by assuming that any choice of $N_{\rho}(\sim N_c)$ vectors out of the possible $N_r(\sim 2N_c)$ is a possible spanning tree,

$$N_{\rm ST} < \binom{2N_c}{N_c} \approx \sqrt{\frac{4N_c}{\pi}} 2^{2N_c}.$$
 (31)

Using this bound we get, in the large N_c limit,

$$\zeta_G < 1.386 + O(\ln N_c/N_c). \tag{32}$$

This bound is too high since most choices of a random set of N_c vectors \vec{r} include forbidden vector combinations of closed loops. A better estimate would be by using a known bound for regular graphs [23,24]:

$$N_{\rm ST}^{(G)} \leqslant \frac{2\ln N_V}{N_V K \ln K} (C_K)^{N_V},\tag{33}$$

where *K* is the valency of the vertices of *G*, and $C_K \equiv [K - 1]^{K-1}/[K(K-2)]^{K/2-1}$. Thus, in the large N_V limit, $\zeta_G \leq \ln C_K$. Almost all the vertices of graphs describing contact network connect four vectors \vec{r} (see, e.g., Fig. 1) and have K = 4. The exceptions are (i) K = 3 for contacts of two-contact particles, but these consist a small fraction, ε , of all particles; (ii) K = 2 for chains of two-contact particles, but these are unstable mechanically and rarer still; (iii) K = 2 for generic boundary contacts, but the number of these scale as $M \ll N_c$. The total number of contacts is $N_c = (N\bar{z} + M)/2$ and the total number of edges is $N_E = N(\bar{z} - \varepsilon)$. Since $M \ll N$ and $\varepsilon \ll 1$ for sufficiently large systems,

$$K = \frac{2N_E}{N_V} \approx 4 \left(1 - \frac{M}{N\bar{z}} - \frac{\varepsilon}{\bar{z}} \right) \approx 4.$$
(34)

Using Eq. (33), we then obtain $C_K = (1.5)^3$ and an improved upper bound, compared to (32),

$$N_{\rm ST} \leqslant (1.5)^{3N_c}, \ \zeta_G \leqslant 1.216.$$
 (35)

However, ζ_G depends not only on *K* but also on further details of the structure. For example, the square and kagome lattices, both regular graphs with K = 4, have different values, $\zeta_{Sq} = 1.166$ and $\zeta_{Kag} = 1.136$ [23,24]. The difference between these structures is their distributions of cell orders, or the *K* value of their dual graphs. The dual of the square lattice is also regular with K = 4, but the dual of the kagome lattice has 2/3 vertices with K = 3 and 1/3 with K = 6. Since, of the two graphs, the latter's distribution of cell orders is closer to that of real systems, in which these range between 3 and 6 (see Fig. 1), then we conjecture that the kagome lattice should describe our systems more closely. This provides us the best estimate,

$$\zeta_G \approx 1.136,\tag{36}$$

which is lower than both Eqs. (32) and (35).

To test this result, we analyzed 2D experimental systems, each of 1172 discs of three different radii, produced by the 3SR Lab, as described in Ref. [25]. We divided the image of each system to several nonoverlapping subsystems of different sizes, and constructed the contact network for each subsystem. Figure 4 demonstrates this procedure and shows one choice of a spanning tree. We counted the number of possible spanning trees for each subsystem using Kirchhoff's theorem [26]. We then calculated ζ_G as a function of system size, using Eq. (30). Figure 5 shows ζ_G as a function of $1/N_c$ alongside ζ_{Sq} and ζ_{Kag} of lattices of the same size, for which the values were also obtained by using Kirchhoff's theorem. Indeed, the value of ζ_G in the disordered systems converges to that of the kagome lattice, supporting our result (36).

Having a reliable estimate of the multiplicity, N_{ST} , we can now estimate the topological entropy:

$$S_{t} = \ln(\Omega_{t}/N_{\rm ST})$$

= 2nN_c ln N_c - (2 ln n! + ζ_{G})N_c + O(n). (37)



FIG. 4. (a) The bottom-left corner of a tridisperse 2D experimental granular system, produced by the 3SR Lab [25]. (b) The network of vectors \vec{r} , circulating every particle and cell. The directionality of the vectors, clockwise around particles and counterclockwise around cells, is not plotted to avoid cluttering. (c) The connectivity network of this system. (d) One choice of a spanning tree, i.e., a subset of independent vectors \vec{r} . All boundary vectors \vec{r} but one are chosen.

In the calculation above, we assumed that $S_a(A)$ depends only weakly on the topology. To test this assumption, we first note that the contribution of the boundary forces to S_a is negligible compared to the structural configurations and that $S_a \approx \ln |B| + O(\sqrt{N})$. Calculating the value of $\ln |B|$ for



FIG. 5. ζ_N , defined in Eq. (30), as a function of the inverse of the number of vertices, $1/N_V$, for different graphs: the kagome lattice (green line), the square lattice (blue line), and the connectivity network of the experimental 2D systems (black open circles, interpolated by red line). The value of ζ_N for the kagome lattice and the experimental systems is calculated numerically, by constructing the graphs and using Kirchhoff's theorem to get the number of spanning trees. ζ_N for the square lattice is calculated analytically. In dashed lines are quadratic extrapolations to $1/N_V \rightarrow 0$.



FIG. 6. The histogram of $\ln |B|$ for 1000 systems of 64 particles each (white bars). Each blue dot represents a single system, at the appropriate *x* value and a random *y* value.

1000 systems of 64 soft disks each, computer-generated by the same protocol [16], we plot its distribution in Fig. 6. Indeed, we find that $\ln |B|$ has a well-defined value with a relative width of 2.9/167 = 1.7%, supporting well the above assumption. Using $N_c = N\bar{z}/2$ and $n = \langle l \rangle/2$, we can express S_t in terms of the number of particles, N:

$$S_t = \frac{\bar{z}}{2} \langle l \rangle N \ln N + \frac{\bar{z}}{2} \left(\langle l \rangle \ln \frac{e\bar{z}}{\langle l \rangle} - \zeta_G \right) N .$$
 (38)

Thus, S_t is dominated by the term $N \ln N$, reflecting the fact that it corresponds to arrangements of all the particles in a network, whose number is of order N!.

This result is very significant: It indicates that the topological structural entropy is extensive, i.e., linear in system size, only when subtracting $(\bar{z}\langle l \rangle/2) \ln N!$ from it. This observation is reminiscent of the much discussed $\ln N!$ subtraction in thermal statistical mechanics. However, it is interesting that, in GSM, only the topological entropy incurs this term. Thus, together with result (17), the total structural entropy, $S_a + S_t$, is extensive. Significantly, relation (38) is supported by, and provides the theoretical explanation for, the numerical observations in Refs. [15,16], which observed directly the N! factor.

VII. DISCUSSION

In this paper, we extended the connectivity-based granular statistical mechanics, proposed in Ref. [18], and derived a number of fundamental results as follows. First, we calculated the complete partition function, Eq. (1), beyond the structural contribution and included the explicit contribution of the stress ensemble. The quadratic exponential in Eq. (7) allowed us to not only solve it, Eq. (10), but also to calculate explicitly expectation values for (i) the connectivity, $\langle C \rangle_a$, showing that there exists an equipartition principle for it; (ii) the squared-norm of the force vector, $\langle \vec{g} \cdot \vec{g} \rangle_a$; (iii) the boundary normal stress, $\langle \sigma_{nn} \rangle_a$; and (iv) the volume, $\langle V \rangle_a$. Using these, we derived *an equation of state*, Eq. (20).

Second, we identified two main sources for the structural entropy, affine and topological, with the former describing microstates with the same topology of the contact network and the latter describing microstates of different topologies. The connectivity-based formulation was shown to be convenient for separating these two contributions, Eq. (27), with all the topology encoded in the connectivity matrix A. This separation made it possible to calculate explicitly each of these contributions. We calculated the affine entropy explicitly from Eq. (16) and found that it scales linearly with the number of particles; i.e., it is extensive.

Third, we established that, to calculate the affine contribution to the partition function, one must take into consideration explicitly the correlations between the DFs as constraints and that ignoring these constraints lead to grossly unphysical results, such as a vanishing mean volume. We then modified the connectivity function to include these constraints, Eq. (24), and showed that these remedy the calculation and give more physical results.

Fourth, we showed that our method of including constraints can be used to describe correlations between the DFs as interactions. This constitutes an extension of the granular statistical mechanics formalism beyond the traditional analog of self-energy-like description. We demonstrated that granular systems possess inherent positive and negative correlations between angles, as well as lengths, of successive DFs along connectivity loops and outlined how to include these as interaction terms in the modified connectivity function. These interactions are convenient in that they resemble nearest neighbor interactions in more traditional systems.

Fifth, we calculated the topological entropy, using the statistics of the matrix A, which describes the topology of a granular assembly as a spanning tree [20]. This calculation was complicated by the concept of *multiplicity*, namely, that a specific structural configuration can be described by N_{ST} spanning trees. We calculated $N_{ST} \sim e^{1.136\overline{z}N/2}$, yielding that the number of microstates is Ω_t / N_{ST} . It follows that the overall entropy is extensive only when subtracting $(\bar{z}\langle l \rangle/2) \ln N!$ from it. This result provides a theoretical explanation of the observations in Refs. [15,16].

Sixth, by calculating explicitly the structure- and forcebased entropies, we established that, in the large system limit, the latter is negligible relative to the former. This result is the direct consequence that the structure phase space is of size N while the force phase space is of size $N^{(d-1)/d}$. Another consequence is that the dependence of expectation values of structural properties on the angoricity tensor [5] is negligible, at least for rigid particles. The implication of this conclusion for force-based expectation values remains to be studied, but we showed that the equation of state (20), which includes the boundary stress, depends negligibly on the angoricity unless its value is extremely small.

Our results hold for an ensemble of mechanically stable granular packs, all generated by the same protocol and all having the same mean coordination number, \bar{z} . Extending the analysis to distributions of \bar{z} across the ensemble is possible, in principle, albeit cumbersome, as it involves enumeration over matrices A of varying dimensions. We reiterating that our estimate of the topological entropy, S_t , was made under the assumption, supported numerically for a certain class of systems in Sec. VI, that different topologies have similar occurrence probabilities. When this assumption does not hold, Eq. (37) must be replaced by the more general Gibbs entropy, $S_t = -\sum P_t \ln P_t$, with P_t the probability of each topology. Thus, Eq. (37) is an upper bound since a uniform distribution maximizes the entropy.

The formalism presented here can be extended readily to 3D systems, following the same conceptual approach. First, one constructs a network of vectors \vec{r} that form a convex hull around each grain [13,19]. Then, choosing a spanning tree of this network, using the same principle as described above, the connectivity function can be calculated straightforwardly. More care is required in calculating the stress ensemble, since 3D cells are surrounded by both grains and throats, which are the openings connecting neighboring cells. Each throat is made of a loop of vectors \vec{r} . As in 2D, the intergranular forces can be solved for in terms of the boundary forces and, if required, additional constitutive relations. The boundary forces are the DFs of the stress ensemble. It then follows that one can define the 3D equivalents of the matrices E and H, and the rest of the analysis is the same as above.

We look forward to numerical and experimental tests of this formulation.

ACKNOWLEDGMENT

S.A. acknowledges financial support by the Alan Howard PhD Scholarship, and thanks S. Martiniani for providing the computer-generated systems [16].

APPENDIX A: CALCULATING EXPECTATION VALUES AND ENTROPY

The different expectation values are all derivatives of $\ln Z_a$ and, using Eq. (10), it is convenient to define

1m 7

$$\ln Z_a = \ln Z_0 + \ln Z_1,$$

(A1)

 $\ln Z_0 \equiv -\frac{d}{2} \ln |B| - \frac{1}{2} \ln |P|_+,$ $\ln Z_1 \equiv \frac{g^2}{2} \operatorname{Tr}(P) + \sum_{\alpha > 0} \ln D(a_i) + \kappa,$ (A2)

and κ is a function of N_{ρ}, M , and g, whose exact form is immaterial at the moment.

(1) The connectivity

The connectivity expectation value is obtained by

$$\langle \mathcal{C} \rangle_a = \tau^2 \frac{\partial}{\partial \tau} (\ln Z_0 + \ln Z_1) \equiv \langle \mathcal{C} \rangle_0 + \langle \mathcal{C} \rangle_1.$$
 (A3)

Starting with $\langle C \rangle_0$, both terms of $\ln Z_0$ are proportional to a power of τ :

$$|B|\sim au^{-N_
ho}, \quad |P|_+\sim au^{d(M-1)},$$

and, using the fact that if $f(x) \sim x^{\alpha}$ then

$$\frac{\partial \ln f}{\partial x} = \frac{\alpha}{x},\tag{A4}$$

we obtain

where

$$\langle \mathcal{C} \rangle_0 = \frac{d\tau}{2} (N_\rho - M + 1). \tag{A5}$$

To calculate $\langle C \rangle_1$, we note that $\ln Z_1$ depends on τ through $\operatorname{Tr}(P) \sim \tau$ and $a_i \sim \sqrt{p_i} \sim \sqrt{\tau}$. One can also obtain from Eq. (11):

$$\frac{\partial \ln D(a_i)}{\partial a_i} = \frac{1}{D(a_i)} - 2a_i, \tag{A6}$$

and thence

$$\langle \mathcal{C} \rangle_1 = \tau^2 \left\{ \frac{g^2}{2\tau} \operatorname{Tr}(P) + \sum_{a_i > 0} \left[\frac{1}{D(a_i)} - 2a_i \right] \frac{a_i}{2\tau} \right\}$$
$$= \frac{\tau}{2} \sum_{a_i > 0} \frac{a_i}{D(a_i)},$$
(A7)

where the first and last terms in the first line cancel out. Summing (A5) and (A7) yields $\langle C \rangle_a$ in Eq. (12).

(2) The volume

The mean volume is obtained from Eq. (18) via

$$\langle V \rangle_a = -2W_{jk} \frac{\partial \ln Z_a}{\partial B_{jk}} + 2U_{jk} \frac{\partial \ln Z_a}{\partial P_{ik}}.$$
 (A8)

Using again (A2), we express a corresponding separation: $\langle V \rangle_a = \langle V \rangle_0 + \langle V \rangle_1$. Starting with $\langle V \rangle_0$ and noting that the dependencies on B_{jk} and P_{jk} originate in |B| and $|P|_+$, respectively, we have

$$\langle V \rangle_0 = -2W_{jk} \frac{\partial \ln Z_0}{\partial |B|} \frac{\partial |B|}{\partial B_{jk}} + 2U_{jk} \frac{\partial \ln Z_0}{\partial |P|_+} \frac{\partial |P|_+}{\partial P_{jk}}$$

Using Eq. (A4) again we have

$$\langle V \rangle_0 = \frac{dW_{jk}}{|B|} \frac{\partial |B|}{\partial B_{jk}} - \frac{U_{jk}}{|P|_+} \frac{\partial |P|_+}{\partial P_{jk}}.$$
 (A9)

To evaluate these expressions, we express the determinants, e.g., |B|, as $|B| = \sum C_{jk} B_{jk}$, which leads to

$$\frac{\partial |B|}{\partial B_{jk}} = C_{jk} = |B|(B^{-1})_{kj}, \tag{A10}$$

with B^{-1} the inverse matrix of *B*. Similarly, for the pseudode-terminant $|P|_+$, we have

$$\frac{\partial |P|_+}{\partial P_{ik}} = |P|_+(\Pi)_{kj},\tag{A11}$$

with Π being the pseudoinverse of *P*. Substituting (A10) and (A11) into (A9) we obtain

$$\langle V \rangle_0 = d \operatorname{Tr}(WB^{-1}) - \operatorname{Tr}(U\Pi).$$
 (A12)

Turning to $\langle V \rangle_1$, ln Z_1 depends on P_{jk} through Tr(P) and the eigenvalues p_i . The former gives $\partial \operatorname{Tr}(P)/\partial P_{jk} = \delta_{jk}$. For the latter, note that if O is the orthogonal diagonalization matrix of P, $(OPO^T)_{ij} = \Lambda_{ij} = p_i \delta_{ij}$, then [27]

$$\frac{\partial p_i}{\partial P_{ik}} = O_{ij}O_{ik},\tag{A13}$$

which also holds for nonzero eigenvalues of a singular matrix.

Together with (A6), these give

$$\langle V \rangle_{1} = g^{2} U_{jk} \delta_{jk} + 2U_{jk} \sum_{a_{i}>0} \left(\frac{1}{D(a_{i})} - 2a_{i} \right) \frac{a_{i} O_{ij} O_{ik}}{2p_{i}}$$

$$= g^{2} \operatorname{Tr}(U) + \sum_{a_{i}>0} \left(\frac{1}{2a_{i} D(a_{i})} - 1 \right) g^{2} U_{ii}'$$

$$= \sum_{a_{i}>0} \frac{g^{2} U_{ii}'}{2a_{i} D(a_{i})},$$
(A14)

where we abbreviated $U_{jk}O_{ij}O_{ik} = (OUO^T)_{ii} \equiv U'_{ii}$. The first and last terms on the second line cancel out because Tr(U) = Tr(U'). Summing (A12) and (A14) gives $\langle V \rangle_a$ in Eq. (19).

(3) *The mean squared norm of the force vector* Noting that the expression

$$\langle \vec{g} \cdot \vec{g} \rangle_a = \frac{1}{Z_a} \int (gg) e^{-\frac{1}{2}\vec{\rho}B\vec{\rho} + \frac{1}{2}\vec{g}P\vec{g}} (d\vec{\rho}) (d\vec{g})$$
(A15)

has the same form as the volume expectation value in Eq. (18), with W = 0 and U = 1, we obtain $\langle \vec{g} \cdot \vec{g} \rangle_a$ by substituting these values in Eq. (19). This gives straightforwardly the result, Eq. (14).

(4) *The entropy*

In analogy to thermal statistical mechanics, the affine entropy is given by

$$S_a = \frac{\langle \mathcal{C} \rangle_a}{\tau} + \ln Z_a. \tag{A16}$$

Its calculation requires the explicit form of κ in Eq. (A2),

$$\kappa = \frac{d}{2} [N_{\rho} \ln(2\pi) + (M-1)\ln(8) + 2\ln(2g)].$$
(A17)

To obtain S_a in Eq. (16), we substitute Eqs. (A2), (12), and (A17) into Eq. (A16).

(5) The boundary normal stress

To obtain an explicit expression for $\langle \sigma_{nn} \rangle_a$, we must calculate it directly from the partition function. We shall use the following calculation:

$$\frac{\int_0^g g' e^{\frac{1}{2}pg^2} dg'}{\int_0^g e^{\frac{1}{2}pg^2} dg'} = \frac{(e^{a^2} - 1)}{\sqrt{2p} e^{a^2} D(a)} = \frac{g(1 - e^{-a^2})}{2aD(a)}, \quad (A18)$$

where $p \ge 0$ and $a \equiv g\sqrt{p/2}$. In the limit $p \to 0$ (A18) reduces to g/2. Using this result, we define

$$\mathcal{G}_j = \frac{1 - e^{-a_j^2}}{2a_j D(a_j)}g\tag{A19}$$

and proceed to calculate the expectation value of the normal component of a boundary force,

$$\langle g_x^i \rangle_a = \frac{1}{Z_a} \int e^{-\frac{1}{2}\vec{\rho}B\vec{\rho}} (d\vec{\rho}) \int g_x^i e^{+\frac{1}{2}\vec{g}P\vec{g}} (d\vec{g}) \,.$$
 (A20)

The first set of integrals cancels out with the corresponding integrals of Z_a . To solve the second set of integrals, we diagonalize P and use (A18) and (A19) to obtain

$$\langle g_x^i \rangle_a = \sum_j O_{ij}^T \frac{\int g_j' e^{\frac{1}{2} p_j g_j'^2} dg_j'}{\int e^{\frac{1}{2} p_j g_j'^2} dg_j'} = \sum_j O_{ij}^T \mathcal{G}_j,$$
 (A21)

with O the diagonalization matrix. Squaring,

$$\langle g_x \rangle_a^2 = \frac{1}{M} \sum_i \langle g_x^i \rangle_a^2 = \frac{1}{M} \sum_{i,j,k} O_{ij}^T O_{ik}^T \mathcal{G}_j \mathcal{G}_k,$$
 (A22)

and using the orthogonality, $\sum_{i} O_{ij}^{T} O_{ik}^{T} = \delta_{jk}$, we get

$$\langle g_x \rangle_a^2 = \frac{1}{M} \sum_j \mathcal{G}_j^2 = \frac{|\mathcal{G}|^2}{M},$$
 (A23)

where $|\mathcal{G}|$ is the norm of a dM-long vector, whose components are \mathcal{G}_j . Combining (A23) with $\langle \sigma_{nn} \rangle_a = M \langle g_x \rangle_a / 4\sqrt{Na}$, as discussed in the main text, gives relation (15).

APPENDIX B: APPROXIMATIONS OF THE DAWSON FUNCTIONS

The partition function contains d(M-1) Dawson integrals, which give rise to cumbersome sums of Dawson functions, $D(a_i)$, in the entropy (16) and the expectation values (12), (14), and (19). In this appendix, we approximate these sums in the limits of large and small $a_i = g\sqrt{p_i/2} \sim g\sqrt{\tau}\gamma_j$, with $\gamma_{j=1,2}$ the two eigenvalues of the inverse angoricity γ . It is convenient to define the dimensionless parameter

$$\chi \equiv g \sqrt{\tau} |\gamma|, \tag{B1}$$

whose limits correspond to the limits of the parameters a_i .

The second-order approximations of $D(\alpha)$, for small and large values of α , are, respectively,

$$D(\alpha \ll 1) = \alpha \left(1 - \frac{2}{3}\alpha^2\right), \tag{B2}$$

$$D(\alpha \gg 1) = \frac{1}{2\alpha} \left(1 + \frac{\ln 2}{\alpha^2} \right).$$
(B3)

In Fig. 7 we show these approximations and observe that they are very accurate, except in the range $0.5 < \alpha < 1.5$, with a relative error of less than 3%. The arguments of the Dawson



FIG. 7. The Dawson function, in solid blue line, alongside its small- and large-parameter two-term approximations, in green and red dashed lines, respectively. The accuracy of the respective approximations is better than 3% outside the range $0.5 < \alpha < 1.5$.

function, a_i , are proportional to the maximal magnitude of a boundary force, g, to $\sqrt{\tau}$, and to the inverse angoricity eigenvalues, $\gamma_{1,2}$. Thus, for a given values of g and τ , the values of a_i s are small at high angoricity and vice versa. Nevertheless, we also need to find a good approximation for the regime $a_i \approx 1$, which we do next.

(1) Approximation of the mean volume

Here, we wish to approximate the sum $\sum 1/[a_i D(a_i)]$ in Eq. (19). From relations (B2) and (B3), we observe that the sum is dominated by the small a_i s. We also note from [25] that, for a typical 2D system and $g\gamma_{1,2}\sqrt{\tau} = 1$ we get $0.5 \leq a_i \leq 200$, with most a_i s smaller than 1. Using then the small-parameter approximation, we have

$$\sum_{a_i>0} \frac{g^2 U'_{ii}}{2a_i D(a_i)} \approx \sum_{a_i>0} \frac{g^2 U'_{ii}}{2a_i^2} \left(1 + \frac{2}{3}a_i^2\right)$$
$$= \sum_{a_i>0} \frac{U'_{ii}}{p_i} + \frac{g^2}{3} \sum_{a_i>0} U'_{ii}$$
$$= \operatorname{Tr}(U\Pi) + \frac{g^2}{3} \operatorname{Tr}(U\Pi P). \quad (B4)$$

For the last step, we used again the diagonalization of Π to establish that the first sum is equal to $\text{Tr}(U'\Lambda^{-1}) = \text{Tr}(U\Pi)$ and the second sum is equal to $\text{Tr}(\Lambda^{-1}\Lambda U') = \text{Tr}(U\Pi P)$. Plugging these into Eq. (19), we obtain for the mean volume

$$\langle V \rangle_a \approx d \operatorname{Tr}(WB^{-1}) + \frac{g^2}{3} \operatorname{Tr}(U\Pi P).$$
 (B5)

(2) Approximation of the mean squared norm of force vector An approximation of expectation value for the boundary forces can be obtained either by following the same route as above or by substituting W = 0 and U = 1 in Eq. (B5), which yields

$$\langle \vec{g} \cdot \vec{g} \rangle_a \approx \frac{g^2}{3} \operatorname{Tr}(\Pi P) = \frac{g^2}{3} d(M-1) .$$
 (B6)

This is again a second-order approximation; to first order, $-\operatorname{Tr}(\Pi)$ of Eq. (14) cancels out with the sum to give, not surprisingly, $\langle \vec{g} \cdot \vec{g} \rangle_a = 0$. This expression is independent of the contactivity, as one would expect intuitively. Its independence of the angoricity is only a feature of the leading term—the next term would be proportional to $\operatorname{Tr}(\Pi PP)$, scaling as $\tau Tr(\gamma^2)$.

(3) Approximations of the mean boundary normal stress and equation of state

For these approximations, we use Eqs. (15), (A19), and (A23). Approximating G_j for small a_j , we have

$$\mathcal{G}_j = \frac{g}{2a_j} \frac{1 - e^{-a_j^2}}{D(a_j)} \approx \frac{g}{2a_j} \frac{a_j^2}{a_j} = \frac{g}{2}.$$
 (B7)

Substituting this in Eq. (15), we get

$$\langle \sigma_{nn} \rangle_a \approx \frac{\sqrt{M}}{4\sqrt{N}a} \sqrt{M\left(\frac{g}{2}\right)^2} \approx \frac{g}{8a}.$$
 (B8)

Using the approximations for $\langle V \rangle_a$, Eq. (B5), and $\langle \sigma_{nn} \rangle_a$, Eq. (B8), we obtain the second-order approximation for the equation of state in the limit of low χ

$$\langle V \rangle_a \langle \sigma_{nn} \rangle_a \approx \frac{g}{8a} \bigg[d \operatorname{Tr}(WB^{-1}) + \frac{g^2}{3} \operatorname{Tr}(U\Pi P) \bigg].$$
 (B9)



FIG. 8. Blue, thin line: the exact affine entropy, calculated according to Eq. (16), for a 2D system of 1172 discs, as a function of the two squared eigenvalues (fixed to be equal) of γ . The contactivity, τ , and the maximal boundary force, g, are set to 1. Green, thick line: the relative error of approximating the Dawson functions, namely the difference between calculating *S* according to Eq. (B11) and according to Eq. (16) over the latter.

(4) Approximation of the entropy

To this end, we need to approximate the expression $\Delta \equiv \sum [\ln D(a_i) + a_i/2D(a_i)]$ in Eq. (16). Using relations (B2) and (B3), for small a_i 's $\Delta \approx 0.5 + \ln a_i$ and for large a_i 's $\Delta \approx a_i^2 - \ln(4a_i)$. It follows that large a_i 's dominate the sum. Moreover, using the definition of a_i , the term $\frac{g^2}{2}$ Tr(P) in Eq. (16) is also proportional to $\sum a_i^2$, making the large a_i limit even more dominant. Using then (B3), we have

$$\Delta = \sum_{a_i > 0} \left[\ln D(a_i) + \frac{a_i}{2D(a_i)} \right]$$

$$\approx \sum_{a_i > 0} \left[a_i^2 - \ln a_i - \ln 4 \right]$$

$$= \frac{g^2}{2} \operatorname{Tr}(P) - \frac{1}{2} \ln |P|_+ - \frac{d}{2} (M - 1) \ln 8g^2. \quad (B10)$$

Substituting this in Eq. (16), we get

$$S_a \approx -\frac{d}{2} \ln |B| - \ln |P|_+ + g^2 \operatorname{Tr}(P) + \frac{d}{2} [N_\rho \ln(2\pi e) + 2\ln(2g) - (M-1)\ln(eg^2)]. \quad (B11)$$



FIG. 9. A numerical check that indeed $\mathcal{G}_j(a_j)$ is bounded between g/2 and g for all j.

The accuracy of this approximation depends on the different noise parameters through the dimensionless parameter χ . Whenever $\chi \gtrsim 1$, the approximation (B11) is accurate to more than 3%. This is demonstrated in Fig. 8 for $\gamma_1 = \gamma_2$ and $\tau = g = 1$ —it shows the relative error, $(S_{approx} - S)/S$, between the approximated entropy, Eq. (B11), and the exact result, Eq. (16), for different values of χ^2 . We find similar graphs when varying τ , g or the ratio of the eigenvalues of γ .

APPENDIX C: ESTIMATION OF THE BOUNDARY NORMAL STRESS

Here, we first establish that $g/2 < \mathcal{G}_j < g$ for all j and, thence, that $|\mathcal{G}| = O(\sqrt{M})$, as stated in Sec. IV. Using the definition of $\mathcal{G}(a_i)$ and the small χ approximation for $D(a_i)$, we have shown in Eq. (B7) that $\mathcal{G}_j(a_j \ll 1) \approx g/2$. Similarly, using the large χ approximation, relation (B3), we have

$$\mathcal{G}_j(a_j \gg 1) \approx g\left(1 - \frac{\ln 2}{a_j^2}\right) < g.$$
 (C1)

This indicates that the value of $\mathcal{G}_j(a_j)$ is bounded between g/2 and g for all j's. We substantiate this by a calculation of the function $D_j(a_j)$, shown in Fig. 9. Thus, $g\sqrt{M}/2 < |\mathcal{G}| < g\sqrt{M}$, and we obtain that, generally, $|\mathcal{G}| = O(M)$.

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