I. INTRODUCTION

Materials composed of hard cohesionless grains, such as dry sand, exhibit many remarkable properties ranging from cluster formation in gaseous phases to unexpected flows and jets in fluid-like phases, and to complex organization of inhomogeneous stresses in dense fluid or solid phases. In this paper we focus on one feature of the solid phase that has received much attention: the distribution \( P(f) \) of contact forces between grains in a system that is supporting a macroscopic compressive stress. In a number of experiments and numerical simulations involving non-cohesive grains, we find that the probability distribution for single-contact forces decays faster than exponentially. This super-exponential decay persists in lattices diluted to the rigidity percolation threshold. On the other hand, for anisotropic imposed stresses, a broader tail emerges in the force distribution, becoming a pure exponential in the limit of infinite lattice size and infinitely strong anisotropy.

We study the uniformly weighted ensemble of force balanced configurations on a triangular network of nontensile contact forces. For periodic boundary conditions corresponding to isotropic compressive stress, we find that the probability distribution for single-contact forces decays faster than exponentially. This super-exponential decay persists in lattices diluted to the rigidity percolation threshold. On the other hand, for anisotropic imposed stresses, a broader tail emerges in the force distribution, becoming a pure exponential in the limit of infinite lattice size and infinitely strong anisotropy.

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In this paper we investigate the question of whether Edwards’ hypothesis leads to exponential tails in \( P(f) \) for a system of non-cohesive grains in which the contact network forms a triangular lattice. We first study the case of hydrostatic compression, where our results confirm those of Snoeijer et al., though we employ different boundary conditions and different numerical methods. We then examine the effects of diluting the lattice to the rigidity percolation threshold. The transition appears to be first order, with a finite jump in the number of available configurations at threshold. \( P(f) \) becomes substantially broader than in the full lattice case, but still decays faster than exponentially on the lattice sizes within our numerical reach. Finally, we consider the effects of anisotropic imposed stresses. We show that strong anisotropy must produce an exponential tail in large systems and present numerical results showing the approach to this limit for varying degrees of anisotropy.

Because the difference between the decay produced by our model and a true exponential decay is rather subtle, it is difficult to give firm interpretations of most of the experimental and numerical results. Observation of two or three decades of exponential decay could still be consistent with the slowly developing deviation we observe in isotropic systems. On the other hand, a given experiment may involve sufficiently strong anisotropies that we would expect something even closer to a true exponential. The recent results of Majmudar and Behringer clearly illustrate that anisotropy due to shearing produces distributions much closer to exponential than those observed under isotropic compression, but we urge caution in making direct comparisons between the model results and experiments on disordered, deformable grains.

In the model described below, all forces are directed along the line of contact between two grains and tensile forces are not allowed. This may be thought of as corresponding to the case of frictionless, non-cohesive,
circular disks, though a generic set of perfectly circular disks could not form as many contacts as are present in the model. Periodic boundary conditions are imposed, so there is no distinction between bulk and boundary contacts. By treating the force network without regard to any distortions of the lattice, we are considering systems of grains with elastic moduli large enough that the boundary forces cause negligible strain; i.e., we consider either very hard grains or very weak boundary forces. In all cases, we fix the contact network and study the ensemble of force configurations on that network. We note that fluctuations are large. \( P(f) \) in any given force configuration may look quite different from the expected distribution. We employ numerical sampling methods involving Monte Carlo moves that maintain the force balance constraints at all times (in contrast to simulated annealing methods).

For the smallest nontrivial lattice, consisting of nine grains, we calculate \( P(f) \) both analytically and numerically, and find good agreement. For larger lattices, we employ numerical sampling methods involving Monte Carlo moves that maintain the force balance constraints at all times (in contrast to simulated annealing methods).

II. THE LATTICE

We study the distribution of bond strengths on an \( n \times n \) triangular lattice corresponding to the contacts in a hexagonal packing of monodisperse circular grains. We consider a system in the shape of a rhombus, having \( n^2 \) bonds or edges in each lattice direction, and subject to periodic boundary conditions, as illustrated in Fig. 1. Each edge carries a scalar variable \( f \) specifying the magnitude of the force transmitted across a contact. In order to fix the three components of the macroscopic stress tensor \( \sigma_{ij} \) we specify the total compressive forces \( F_1, F_2, \) and \( F_3 \) supported along each of the three lattice directions. There are \( 3n^2 \) variables in the system. For each grain, the vector force balance condition imposes 2 constraints. These constraints are not all independent, however. The periodic boundary conditions guarantee that the sum of all the single-grain constraint equations is trivially zero, leaving \( 2n^2 - 2 \) independent constraints. Fixing \( F_1, F_2, \) and \( F_3 \) imposes 3 additional constraints, for a total of \( 2n^2 + 3 \) constraints. This leaves \( n^2 - 1 \) degrees of freedom in the force configuration.

In addition to force balance equations, there are inequality constraints associated with the fact that the material is non-cohesive. No force \( f \) is allowed to be negative. It is this condition that introduces nonlinearity in the system. For any two force balanced, tension free configurations on the lattice, any weighted average with positive weights will also be force balanced and tension free. Sums with a negative weighting of a configuration, however, will not always be allowed, as they may contain negative forces on some edges.

We represent the system configuration by a vector of forces \( f_i, i = 1 \ldots 3n^2 \). The above constraints create a space of possible configurations that fills a finite, convex \( n^2 - 1 \)-dimensional volume, with boundaries determined by the inequalities \( f_i \geq 0 \). Our interpretation of the Edwards hypothesis is to assign a uniform probability (Lebesgue) measure on the space of allowed configurations.

III. ISOTROPIC LATTICE

In principle, \( P(f) \) can be calculated in the following way. Fix one edge to carry force \( f_i \); this restricts the system to a \( (n^2 - 2) \)-dimensional subset of the \( (n^2 - 1) \)-dimensional allowed volume \( V \) in configuration space. \( P(f) \) is simply the ratio of the \( (n^2 - 2) \)-dimensional “area” to the \( (n^2 - 1) \)-dimensional total volume. More formally, let \( E_m \) be the set of 6 edges touching node \( m \); let \( e_g^{(m)} \), \( g = 1 \ldots 6 \), be a unit vector along edge \( g \) in \( E_m \) pointing towards node \( m \); and let \( L_k, k = 1 \ldots 3 \), be the set of \( n^2 \) edges along lattice direction \( k \). Then we have

\[
P(f) = \frac{1}{V} \left[ \prod_{j=1}^{3n^2} df_j \right] \delta \left( f_i - f \right) \prod_{m=1}^{n^2} \delta \left( \sum_{f_g \in E_m} f_g e_g^{(m)} \right) \prod_{k=1}^{3} \delta \left( nF_k - \sum_{f_l \in L_k} f_l \right)
\]

(1)
where edge $i$ has been fixed to have value $f$. The integral is taken over all $f$’s. The first delta function ensures that edge $i$ carries force $f$. The next ensures force balance at each vertex, and the last enforces the boundary conditions. The integral represents the volume of a $(n^2 - 2)$-dimensional slice of a polytope, therefore $P(f)$ on an $n \times n$ lattice is a piecewise polynomial of order $n^2 - 2$.

The average force on the edges in $L_k$ is $F_k/n$. The last factor in Eqn. (1) requires comment. For definiteness, consider $k = 2$. Let us divide the set of edges in $L_2$ into $n$ layers, each layer containing the $n$ edges that intersect a line along one of the other edge directions. The following argument shows that the sum of the $f_j$ on each layer must be equal to $F_2$. Fig. 2 shows a shaded slab of the system. The total force on this slab must be zero, which implies that the vector sum of the forces on the edges indicated by thick lines on one side of the slab must be equal to the vector sum of the forces on the other side. Since the vector sum has a unique decomposition into contributions from the $L_2$ edges and the $L_3$ edges, the sum of the $f_j$’s in each direction must independently be the same on both sides of the slab. This argument is independent of the thickness of the slab, so in the $L_2$ direction each of the $n$ layers must have $f_j$’s that sum to $F_2$, and similarly for $L_1$ and $L_3$.

For the $3 \times 3$ case, the integral can be evaluated analytically. Here we state the isotropic result: the anisotropic result is presented below. The calculation is detailed in Appendix A. (We have not found an analytic expression for $P(f)$ for arbitrary lattice size. See Ref. [10], however, for a treatment of the case in which the sum $F_1 + F_2 + F_3$ is fixed, but not the individual $F_k$.) In the isotropic case ($F_1 = F_2 = F_3 = F$), we find

$$P(f) = \frac{8}{45F^5} \Theta(f)\Theta(F - f)(F - f)^2$$

$$\times (5F^5 + 73fF^4 - 111f^2F^3 + 125f^3F^2 - 59f^4F + 9f^5)$$

where $\Theta(f)$ is the Heaviside step function. This expression is plotted in Fig. 3.

On larger lattices we employ numerical methods to measure $P(f)$. This requires generating numerous configurations in the allowed volume of configuration space in a manner consistent with the Edwards flat measure. Previous work has implemented a simulated annealing algorithm, which generates each configuration by starting from a random point in the space of all possible $f_i$ and relaxing to some point in the lower dimensional subset of interest. We employ a different technique in which, starting from one force-balanced configuration – a point in the allowed subset of stress states – new configurations are generated via moves that always produce allowed configurations. By identifying a set of moves that span this compact space, reach any set of positive volume in a finite time, and involve symmetric transition probabilities, we can be sure that at sufficiently long times the space is being sampled with uniform measure. These moves are described below.

For the $n \times n$ lattice with $n \geq 3$, we construct a set of $n^2$ wheel moves, one centered on each node. The wheel move associated with a given node acts on the nearest neighbor edges (“spokes”) and next nearest neighbor edges (“rim”) of the node, reducing one set of $f$’s and augmenting the other by the same amount. The move is implemented in the following steps. Identify the smallest force on the spokes and call it $s_{\min}$; likewise, call the smallest force on the rim $r_{\min}$. Randomly choose a force increment $\Delta f$ with uniform measure on the interval $[-s_{\min}, r_{\min}]$. Adding $\Delta f$ to every edge on the spokes and subtracting it from every edge on the rim, which respects force balance on every node touching the wheel, constitutes a wheel move.

In the space of linear combinations of wheel moves, there is one obvious null direction; i.e., a linear combination of wheel moves that results in no change to any edge of the lattice. We prove in Appendix B that the number of linearly independent wheel moves is exactly $n^2 - 1$, which is the number required to span the space of force configurations.

Fig. 4 illustrates a geometric subtlety that must be taken into account when attempting to explore the al-
lowed stress states using wheel moves (or any other basis set of moves). Since the given pair of basis vectors span the two-dimensional convex space shown, it is clearly possible to navigate from any interior point to any other by making discrete steps along the basis vectors. It may not be possible, however, to make a move in either basis direction if the initial configuration lies exactly on a corner. In higher dimensions it is likewise possible to be stuck on a corner or a boundary of higher dimension. It is therefore important to find initial configurations that lie in the interior of the space, rather than on a boundary, for the purposes of our Monte Carlo sampling technique. An algorithm such as the one described above, beginning at an interior point, can come arbitrarily close to all boundaries but will never reach them. Regions near corners are visited infrequently but for long times in such a way that, for sufficiently long runtime, the space is sampled with uniform measure. 

Fig. 5 shows the agreement between the exact calculation and simulation on the $3 \times 3$ lattice. The forces are normalized by choosing $F$ such that the average force $\langle f \rangle$ is unity. The peak near $\langle f \rangle$ is typical for larger lattices as well, but finite size effects are clearly evident in the tail, since $P(f)$ must go to zero at $f = F$ (here $F = 3$).

A typical configuration for the $15 \times 15$ lattice is shown in Fig. 6. $P(f)$'s for the cases $n = 5, 10, 15$, and 20 are shown in Fig. 7. The peak near $\langle f \rangle$ is again apparent, and for small $f$ the curves appear to coincide, though small differences exist that are masked by the logarithmic scale.

For $f \gtrsim 3\langle f \rangle$ the curves separate, with the $n = 5$ distribution decaying most rapidly. The four distributions in the figure all decay faster than exponentially, and they broaden slightly with increasing $n$.

To gain confidence that the curves are converging to a large system limit, we measured correlations between forces as a function of the distance between edges. In directions both longitudinal and transverse to a particular edge, as shown in Fig. 8, we see that on a $20 \times 20$ lattice correlations have decayed to the 1% level at a distance of 10 lattice constants. On larger lattices we see exponential decay with a decay length of approximately 3 lattice constants, although much more data on larger lattices would be necessary to measure this precisely. We therefore expect to see little difference in $P(f)$ for lattices larger than $n = 20$. Supporting this expectation, there is very little difference between the $n = 15$ and $n = 20$ distributions for $f$ up to the largest values we have measured, which covers five decades of $P$. Though it is conceivable that the asymptotic form of $P(f)$ at large $f$ in our model is exponential, the domain displayed in Fig. 7 is the relevant one for comparison with experiments, and it clearly
FIG. 8: Above, correlations are calculated in the longitudinal and transverse directions. Below, correlation function \( g(r) = \langle f_i f_{i+r} \rangle - \langle f_i \rangle^2 \) on a 20 × 20 lattice. The correlation is computed for edges that have the same orientation. The longitudinal correlation refers to displacements by \( r \) lattice constants along the direction of the edge. The transverse correlation refers to displacements perpendicular to the edge. The plots above are averaged over multiple reference edges. The transverse correlations are in fact negative, but we take their absolute value to facilitate plotting on a log scale.

shows a decay that is faster than exponential.

### IV. DILUTED LATTICES

Real disks are not perfectly monodisperse, so in any hexagonal packing of hard disks some of the bonds on the triangular lattice will not actually be present. For a given stress state, a diluted lattice has fewer bonds to carry the same force. Those remaining will, on average, carry more force, resulting in a shift in \( P(f) \) toward larger forces, which may involve a broadening of the tail. As the number of edges removed from the lattice increases, the force configurations tend to become less homogeneous, with strong forces concentrated in chain-like structures. As more edges are deleted there comes a point where the lattice can no longer support the imposed stress; this is the rigidity-percolation transition. Fig. 9 shows a typical force configuration for a randomly diluted lattice. We study randomly diluted lattices at the transition to see whether the broadening takes the form of an exponential decay.

The random dilution process merits further remark. For a random process in which each bond is removed with probability \( \phi \), the infinite triangular lattice under isotropic compression cannot support stress for any \( \phi > 0 \). Thus in an \( n \times n \) simulation the fraction (not the number \( n_d \)) of deleted edges at the rigidity percolation threshold goes to zero as \( n \to \infty \). As an arbitrarily large real hexagonal packing does support compression, this means that the process of bond breaking in the real packing does not happen randomly but rather in a correlated way. Nevertheless, our simulations employ random dilution on finite lattices. We also neglect for present purposes the possible buckling instability of a force chain. Disallowing configurations that might buckle could only result in ensembles with fewer dilutions and therefore would not change the conclusions described below.

We construct lattices at the rigidity percolation threshold in the following way. Given the imposed macroscopic stress and beginning with an empty lattice, edges are selected at random to be added to the lattice. After each addition, we check to see if the lattice is capable of supporting the imposed stress using the simplex algorithm. We take the lattice to be at threshold when it first supports the imposed stress, at which point we know that there exists at least one edge such that, when removed, the system could no longer support the stress. The edges that have not been added to the lattice at this stage are called “deleted edges.” Our construction yields a lattice in which deletions are randomly distributed in space. Note that the process will in general yield some edges that are present but not connected to the edge network in a way that allows them to bear any force. We refer to these edges as “effectively deleted.”

These effectively deleted edges on a lattice supporting compressive forces can be identified in the following way. Consider the 6 edges that meet at a given node. An edge becomes effectively deleted if its opposite edge has been deleted (possibly effectively) and at least one of the edges making a 120° angle with it has also been deleted (possi-
bly effectively). Under these conditions any force on the edge in question can not be balanced by positive forces on the other edges sharing the node. To find all of the effectively deleted edges we examine all nodes repeatedly until no new deletions are found.

The wheel moves used for investigation of the undiluted lattice no longer work on the diluted lattice. The wheel moves add or subtract a quantity $\Delta f$ to each edge they touch, but this is impossible if one of the edges is deleted. Therefore each deleted edge renders unusable the four wheel moves to which it belongs. To salvage a deleted edge renders unusable the four wheel moves to which it belongs. To salvage a deleted edge requires an effective deletion of the edge that touches which preserve the deleted edge unaffected. For a single deleted edge there are three linearly independent combinations of the wheel moves that span the appropriate space, linear combinations of the wheel moves can be formed that leave the deleted edge unaffected. For a single deleted edge there are three linearly independent combinations of the wheel moves that leave the deleted edge unaffected. For a single deleted edge there are three linearly independent combinations of the wheel moves that leave the deleted edge unaffected. For a single deleted edge there are three linearly independent combinations of the wheel moves that leave the deleted edge unaffected. For a single deleted edge there are three linearly independent combinations of the wheel moves that leave the deleted edge unaffected. For a single deleted edge there are three linearly independent combinations of the wheel moves that leave the deleted edge unaffected.

We refer to a particular linear combination of wheel moves that remains a viable move on a diluted lattice as a “multi-wheel move.” Let the number of linearly independent multi-wheel moves be $N_m$. Just as the undiluted lattice had $n^2$ wheel moves and $n^2 - 1$ degrees of freedom, the diluted lattice having $N_m$ multi-wheel moves has $N_m - 1$ degrees of freedom. Finding a complete set of multi-wheel moves then provides not only a means to perform numerical simulation but also a count of the degrees of freedom in the diluted system.

A linearly independent set of $N_m$ multi-wheel moves on the diluted lattice can be constructed as follows. For $j = 1 \ldots n^2$, form a vector $\mathbf{\vec{w}}_j \in \mathbb{R}^{n^2}$ such that the components of $\mathbf{\vec{w}}_j$ specify the effect of the wheel move centered on node $j$ on the deleted edges. That is, $(\mathbf{\vec{w}}_j)_i = +1$ if the $i^{th}$ deleted edge is a spoke of node $j$, $-1$ if the $i^{th}$ deleted edge is a rim of node $j$, and 0 otherwise. Form the $n^2 \times n^2$ matrix $W_{ij} = (\mathbf{\vec{w}}_j)_i$. Consider a linear combination of wheel moves with coefficients $m_k, k = 1 \ldots n^2$. This linear combination has no effect on deleted edges if and only if $W \mathbf{\vec{w}} = 0$. Thus $N_m = \dim(N(W) = n^2 - \text{rank}(W)$, where $N(W)$ is the null space of $W$.

Note that we demand the wheel moves respect both real and effective deletions. Were we not to specify effective deletions as well, the algorithm would in general produce additional multi-wheel moves which were nonzero on some effectively deleted edges. This is because the multi-wheel moves could also act on a lattice that supports tensile as well as compressive forces. On such a lattice there are fewer effective deletions; in particular, a node could be force-balanced having only three edges all on the same side of a line through the node. To model a noncohesive material, then, we must include these additional effective deletions as well.

We use the following technique to investigate the degrees of freedom in lattices at threshold. Edges are added to a lattice one at a time until the rigidity percolation threshold is reached, the set of wheel moves on that threshold lattice are constructed to determine the dimension of the volume of allowed configurations in state space $N_m - 1$, and the process is repeated for a number of threshold lattices. We refer to the system as having $N_m - 1$ degrees of freedom, as this is the number of multi-wheel moves it possesses. These lattices have differing numbers of deleted edges. For lattice with the same number of deleted edges $n_d$ we calculate the average ratio of the number of degrees of freedom of the dilute system to the number in an undiluted one: $\langle \delta \rangle = ((N_m) - 1)/(n^2 - 1)$. For each plot a number of lattices at the rigidity-percolation threshold were generated and the degrees of freedom and deleted edges counted. For large enough lattices, the system has a large number of degrees of freedom available as soon as rigidity-percolation is reached. Note that the observed interval of $\phi$ values shifts as the lattice size is increased.

![Figure 10: The ratio of average number of degrees of freedom in threshold lattices to the number in the undiluted case.](image)

FIG. 10: The ratio of average number of degrees of freedom in threshold lattices to the number in the undiluted case $\langle \delta \rangle = ((N_m) - 1)/(n^2 - 1)$ vs. the ratio of deleted edges in the threshold lattice to the total number of edges $\phi = n_d/3n^2$. A discrete jump in $\langle \delta \rangle$ emerges as $\phi$ is decreased for systems larger than $15 \times 15$, indicating that the transition is first order. This suggests that $P(f)$ near the transition point is unlikely to show qualitatively different behavior from that of the undiluted lattice, though a quantitative broadening of the force distribution is expected.

We measure $P(f)$ averaged over a number of threshold $20 \times 20$ lattices. The resulting distribution is indeed much broader than the undiluted case, but there is still curvature in the distribution on a linear-log plot. Numerical studies in this regime are hampered by two factors. First, each multi-wheel move requires the examination of many edges to ensure that no $f_j$ becomes negative. Second, the maximum size of a multi-wheel move is typically
to stresses imposed by choosing different initial configurations and run 2.
In the second case, we consider a single lattice at threshold however,
results are shown in Fig. 11. As may be expected given the large
percolation threshold does not produce exponential tails.

We have gathered data using two procedures. In one case we construct
a large number of threshold lattices, find initial configurations
we use the simplex method several times with different coefficients and average
data together to determine samples with uniform measure is still convex but contains
many corners that are not easily accessed. The data obtained for a given lattice may exhibit a
bias depending on the initial configuration unless the number of moves considered is extremely large.

In the second case, we consider a single lattice at threshold (with a typical number of
degrees of freedom), find 25 different initial configurations and run $2.5 \times 10^6$ multi-wheel moves from each one, averaging the force data to obtain
the $P(f)$ associated with that particular lattice. The results are shown in Fig. 11. As may be expected given the first-order nature of the rigidity percolation transition, neither procedure produces an exponential tail in $P$ at large $f$. We conclude that random dilution to the rigidity percolation threshold does not produce exponential tails for triangular lattices.

V. ANISOTROPIC LATTICE

We now consider the undiluted lattice with anisotropic stresses imposed by choosing $F_1$ different from $F_2$ and $F_3$. For example, one lattice direction may be subject to stronger compression than the others, creating qualitatively distinct force distributions in the strong and weak directions. We will show that in strongly anisotropic systems the strong direction contributes an exponential decay to the tail of $P(f)$.

We consider $F_1 = F + \Delta$, $F_2 = F_3 = F$, and parameterize the anisotropy by $\alpha = (F + \Delta)/F$, so that $\alpha = 1$ corresponds to isotropic stress. In terms of $\langle f \rangle$, $F = 3\alpha \langle f \rangle/(2 + \alpha)$. The average force in the weak direction $\langle f_w \rangle = F/n$, while that in the strong direction is $\langle f_s \rangle = (F + \Delta)/n$. Similarly, forces in the weak direction can not exceed $F$, and those in the strong direction can not exceed $F + \Delta$. Forces in the strong direction will populate the tail of $P(f)$.

In the limit $\alpha \to \infty$ the system becomes much simpler: only the strong direction carries force, doing so via $n$ chains of $n$ edges, each edge in a given chain carrying the same force as shown in Fig. 12. The forces on the $n$ chains must sum to $\Delta$.

Simple dimensional analysis of the scaling of the allowed volume when one chain is fixed at force $f$ yields

$$\lim_{n \to \infty} P(f) \propto \lim_{n \to \infty} \left(1 - \frac{f}{n \langle f_s \rangle}\right)^{-2} = e^{f/(\langle f_s \rangle)}$$

We see that in this limit anisotropy induces an exponential tail in $P(f)$. We next investigate the extent to which the limiting behavior is reflected at finite anisotropies.

To gain some intuition for the approach to the anisotropic limit we return to the $3 \times 3$ case, which can be solved analytically. We let $P_s(f)$ and $P_w(f)$ denote the separate distributions of forces in the strong and weak directions, respectively. $P_w(f)$ is identical to $P(f)$ in Eqn. 8. In the strong direction, however, the situation is more complicated. A complete expression is given in Appendix C.

For the $3 \times 3$ case, the fully anisotropic limit $\alpha \to \infty$ has $P(f) = \delta(f_s) - 2(3f_s - f)$, $f \in [0, 3f_s)$, a piecewise linear function. For $\alpha > 2$, $P(f)$ is given by $P_s^{(2\theta)}(f)$ (from Appendix C) between $F$ and $\Delta$. $P_s^{(2\theta)}$ is a linear function, and the interval from $F$ to $\Delta$ grows with $\alpha$. As $\alpha$ increases further the width of this region grows and the slope approaches the limiting slope of $-\delta(f_s)^{-2}$. $P_s(f)$ is plotted for increasing $\alpha$ in Fig. 13.
FIG. 13: $3 \times 3$ triangular lattice under anisotropic stress. The force distribution $P_s(f)$ for forces in the strong direction is plotted for increasing anisotropy. Each distribution is scaled to $\langle f_s \rangle = 1$. The limiting form is a piecewise linear function of $f$. $P_s$ for finite anisotropy has a linear region in the middle; this region grows in width and approaches the limiting slope as anisotropy is increased.

FIG. 14: A $15 \times 15$ anisotropic lattice with $\alpha = 5$. The force on an edge is redundantly mapped to color and width (stronger forces are darker and thicker). The stronger stress is imposed in the horizontal direction.

On larger lattices, numerical calculations show similar behavior to the $3 \times 3$ case, but with the limiting linear distribution replaced by the limiting exponential of the large system limit over the range of forces of interest. (The distribution must be cut off at $f = F_s$.) Fig. 15 shows a typical anisotropic configuration. Fig. 15 shows $P(f)$ for anisotropic lattice of size $15 \times 15$, with the contributions from the weak and strong directions shown separately. The role of the strong direction in broadening the distribution is clear. Results are shown for two cases, one in which strong forces are imposed on a single lattice direction and another in which strong forces are imposed on two lattice directions. Note the difference in behavior for small $f$. In the latter case there is no peak in $P(f)$ for small $f$.

Fig. 16 shows force distributions on a $15 \times 15$ lattice for several values of $\alpha$. For sufficiently large $\alpha$ a middle portion of $P(f)$ appears to be exponential. As $\alpha$ increases, so does the extent of this portion. We hold $\langle f_s \rangle$ fixed so that every curve will have the same decay length.

We conclude that $P(f)$ on the anisotropic lattice displays an exponential tail in the following sense. For moderate $\alpha$ and $n$, a portion of the tail of $P(f)$ is nearly linear on a log plot. The decay is not truly exponential because of the finite anisotropy and finite lattice size, but the limiting behavior of $P(f)$ as $n$ and $\alpha$ are increased is a true exponential. The region of nearly exponential decay grows with increasing anisotropy, extending as far as $f \approx 3\langle f_s \rangle$ for $\alpha = 8$.

We note that the degree of anisotropy represented by a given value of $\alpha$ can be compared to the anistropy supported by materials described by an internal friction parameter. The ratio of major and minor principal stresses on our lattice is $\sigma_1/\sigma_2 = (\alpha + 1)/\sqrt{3}$. By exploiting the relation $\sigma_1/\sigma_2 = (1 + \sin \phi)/(1 - \sin \phi)$ [30], where $\phi$ is the angle of internal friction, we find that $\alpha = 8$ corresponds...
to an internal friction of approximately 43°.

The survival of the exponential tail of strong forces in large triangular lattices under anisotropic loading can be traced to the discussion based on Fig. 12 concerning the sums of forces along layers of edges in the various directions. That argument shows that the strong forces in one lattice direction cannot be redistributed into the other directions due to vector force balance constraints. Thus for strongly anisotropic lattices, the strong forces will follow scaling laws close to the limiting exponential form corresponding to no force at all in the weak direction(s). Note that in the case of two strong directions and one weak, the two strong are effectively independent since strong chains from the two directions cannot interact significantly without generating a strong chain in the supposedly weak direction and thereby reducing the degree of anisotropy.

VI. CONCLUSION

We have investigated the distribution $P(f)$ of contact forces on a lattice of triangular bonds under the Edwards flat measure. An interesting question is whether one expects exponential decay in $P(f)$. The triangular lattice is a simple but nontrivial system for studying this phenomenon.

The distribution on the $n \times n$ periodic lattice decays faster than exponentially, as reported previously. Diluting the lattice induces significant broadening in $P(f)$, but the decay remains faster than exponential. Even at rigidity percolation associated with random bond dilution of the lattice, no qualitative change in the form of $P(f)$ is discernible. In particular, we do not see evidence for an exponential tail associated with the transition. Consistent with these direct measurements of $P(f)$, we find that the transition is first order, which generally suggests that no qualitative changes should be expected as the percolation threshold is approached.

On the other hand, imposing anisotropic stress on the undiluted lattice can induce an exponential tail in $P(f)$. In the limit of an infinite lattice with stress imposed only along one lattice direction, the distribution of contact forces is a pure exponential. Numerical simulation shows that evidence of this behavior may still be seen for finite lattice sizes and a finite ratio of the compressive forces in the strong and weak directions. In such a scenario the tail of $P(f)$ is not a true exponential, but appears approximately linear on a log plot of $P(f)$ for some interval in $f$. For large enough anisotropies, $P(f)$ decays three orders of magnitude from its maximum and $f \approx 3(f_c)$ before deviation from exponential decay becomes obvious.

In the triangular lattice model, anisotropy is associated with the appearance of long force chains oriented along the strong direction, a phenomenon that has also been observed in experiments on disordered systems. Extension of the numerical techniques employed above to rigid bars that form a disordered triangulation of the plane would be straightforward. Wheel moves, for example, would be defined by taking all bars sharing a given vertex to be the spokes, though it would no longer be true that all bars receive force increments of the same magnitude during a move. The primary difference between the perfect triangular lattice and disordered ones (or even other crystalline ones) is that lattices with no sets of collinear bars that span the system cannot support arbitrarily strong anisotropic stresses; i.e., in most cases there would be a maximal value of $\alpha$. A detailed investigation of the effects of disorder would be of interest.

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APPENDIX A: ANALYTIC $P(f)$

$P(f)$ for the $3 \times 3$ isotropic lattice is calculated as follows. There are $3^2 - 1 = 8$ degrees of freedom; we take nine basis elements subject to a constraint which is to be imposed by hand. Three of these elements are shown in Fig. 17. There are three honeycomb elements, $\phi_i$, $i = 1 \ldots 3$, three elements $\psi_{j1}$, $j = 1 \ldots 3$, and three elements $\psi_{j2}$. The figure depicts $\phi_i$, $\psi_{11}$, and $\psi_{12}$. The $\psi$ elements make no net contribution to the total force in the system. The three $\phi$ elements must sum to $F$; this is the additional constraint we impose. All elements are isotropic by construction.
The wheel move with coefficients \( \{m_i\} \) has no effect on the given node if and only if
\[
m_i + m_j - m_k - m_l = 0. \tag{B1}
\]
To interpret Eqn. [B1] geometrically, let \((\mu_i, \nu_i)\) be the coordinates of node \(i\) in Fig. 13a, and define \(X_i \in \mathbb{R}^3\) by
\[
X_i = (\mu_i, \nu_i, m_i). \tag{B2}
\]
Define \(X_j, X_k,\) and \(X_l\) similarly. Then Eqn. [B1] is satisfied if and only if \(X_i\) lies in the plane in \(\mathbb{R}^3\) determined by \(X_j, X_k,\) and \(X_l\).

With further application of Eqn. [B1] we can extend the conclusion to all vertices of the outer triangle, and this may be continued to conclude that all points \(X_i\) lie in a single plane. Finally, periodicity requires that this plane is horizontal: i.e., \(m_i = m_j \) for all \(i\) and \(j\). QED

### APPENDIX C: ANISOTROPIC \(P(f)\) ON THE \(3 \times 3\) LATTICE

The calculation of \(P(f)\) for the anisotropic \(3 \times 3\) case proceeds similarly to the isotropic calculation. The anisotropy requires the division of the relevant integral according to whether the force \(f\) is greater than or less than the difference \(\Delta\) between the strong \(F_i\) and the weak ones. For the case \(\Delta < F\), we find
\[
P_s(f) = \frac{1}{N} \left\{ \begin{array}{ll}
P_s^{(1)}(f) & f < \Delta \\
P_s^{(2a)}(f) & \Delta < f < F \\
P_s^{(3)}(f) & F < f < F + \Delta \\ \end{array} \right.
\tag{C1}
\]
where \(V\) is a normalization constant and the \(P_s\)’s are given below. For the case \(\Delta > F\)
\[
P_s(f) = \frac{1}{N} \left\{ \begin{array}{ll}
P_s^{(1)}(f) & f < F \\
P_s^{(2b)}(f) & F < f < \Delta \\
P_s^{(3)}(f) & \Delta < f < F + \Delta. \\ \end{array} \right.
\tag{C2}
\]
The functions \(P_s\) in Eqn. [C2] are

\[
P_s^{(1)}(f) = (7\alpha - 2)F^7 + 21(4\alpha - 1)F^6f - 42(5\alpha + 1)F^5f^2 + 140(2\alpha + 1)F^4f^3 - 210(\alpha + 1)F^3f^4 + 84(\alpha + 2)F^2f^5 - 14(\alpha + 5)Ff^6 + 12f^7; \tag{C3}
\]
\[
P_s^{(2a)}(f) = (3\alpha^2 - 14\alpha^2 + 21\alpha^2 - 35\alpha^2 + 42\alpha^2 - 14\alpha + 2)F^7 + 21(\alpha^2 - 4\alpha^2 + 5\alpha^2 - 5)\alpha^2F^6 f + 21(3\alpha^4 - 10\alpha^4 + 10\alpha^2 - 15)\alpha^5F^5 f^2 - 35(3\alpha^4 - 8\alpha^4 + 6\alpha^2 - 8\alpha - 5)F^4f^3.
\]
\[ +105(\alpha^3 - 2\alpha^2 - \alpha - 2)F^3f^4 - 21(3\alpha^2 - 8\alpha - 7)F^2f^5 + 7(\alpha - 12)Ff^6 + 9f^7; \]  
\[ P_s^{(2\alpha)}(f) = (21\alpha - 4)F^7 - 21F^6f; \]  
\[ P_s^{(3\alpha)}(f) = (\alpha F - f)^2 \left[-3\alpha^5 - 14\alpha^4 - 21\alpha^3 + 35\alpha - 42\right]F^5 + (15\alpha^4 - 56\alpha^3 + 63\alpha^2 - 35)F^4f \] 
\[ + (30\alpha^2 - 56\alpha + 21)F^2f^3 - 3(10\alpha^2 - 28\alpha + 21)\alpha F^3f^2 - (15\alpha - 14)Ff^4 + 3f^5 \right]. \] 

\[ (\text{C}4) \]

\[ (\text{C}5) \]

\[ (\text{C}6) \]