Evolution of Force Networks in Dense Particulate Media
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We introduce novel sets of measures with the goal of describing dynamical properties of force networks in dense particulate systems. The presented approach is based on persistent homology and allows for extracting precise, quantitative measures that describe the evolution of geometric features of the interparticle forces, without necessarily considering the details related to individual contacts between particles. The networks considered emerge from discrete element simulations of two dimensional particulate systems consisting of compressible frictional circular disks. We quantify the evolution of the networks for slowly compressed systems undergoing jamming transition. The main findings include uncovering significant but localized changes of force networks for unjammed systems, global (system-wide) changes as the systems evolve through jamming, to be followed by significantly less dramatic evolution for the jammed states. We consider both connected components, related in loose sense to force chains, and loops, and find that both measures provide a significant insight into the evolution of force networks. In addition to normal, we consider also tangential forces between the particles and find that they evolve in the consistent manner. Consideration of both frictional and frictionless systems leads us to the conclusion that friction plays a significant role in determining the dynamical properties of the considered networks. We find that the proposed approach describes the considered networks in a precise yet tractable manner, allowing to identify novel features which could be difficult or impossible to describe using other approaches.

PACS numbers: 45.70Qj, 83.80Fg

I. INTRODUCTION

Particulate systems have been extensively studied through centuries due to their importance to our everyday life. These systems appear everywhere, from nano to cosmic scales, and may evolve either hard particles (such as sand) or soft ones (emulsions, foams), see, e.g. [1–4]. As these system are exposed to some external (e.g. compression) or internal (electric, magnetic, gravitational...) influence they may compress, reaching a stage that the particles are in more-or-less permanent contacts. As the systems evolve in time (for whatever reason), the contacts between them in general evolve as well.

Formation of contact networks between the particles, and their properties, have been extensively studied in many different contexts, and using a number of different tools including percolation and network type of approaches; see [5, 6] for overviews. In addition to contact networks, however, there is an additional network of interactions, often called force network in the literature related to granular materials. These force networks are not simply slaved to contact networks, due to indeterminacy of the interaction between particles. A simple example of this is the fact that multiple force networks may be consistent with the condition of force and torque balance in a system. These force networks include complete information about a system and therefore is of significant interest to describe and eventually understand their properties, in particular since it is well known that the interparticle forces play a key role in determining the mechanical properties of static and dynamic systems; see e.g. [6] for an extensive review in the context of amorphous solids.

Physical systems of relevance typically consist of large number of particles, and therefore the force networks may become extremely complex. Due to this complexity, it is necessary to develop techniques that lead to an understanding of the important properties of these networks, without necessarily considering all the details, since this would lead to an intractable study. One obvious idea is to consider statistical properties of these networks, and to ask, e.g., what is the probability of having a force between two particles of a given magnitude. Such studies have been carried out for granular and other systems (see, e.g., [4, 7, 8]) and have led to a significant new insight. Even on the statistical level, however, there are still open questions - one example is a recent discussion of the probability of presence of large forces in granular...
systems \cite{1,9}.

Going beyond purely statistical description requires analyzing in more detail the local properties of the force networks. Such studies have been extensively utilized only recently, but have already indicated the complexity of the problem. Examples include detailed discussion of the forces between particles, see \cite{10,12} and the references therein, where these local properties were connected to global response of the systems considered. Going beyond statistical level, and in the same time attempting to keep the focus on the global structure of the interaction networks requires implementation of new techniques. Considered approaches include network-type of analysis \cite{13,15}. These works provide a significant new insight and confirm that the properties of force networks are relevant in the context of propagation of acoustic signals \cite{13}, fracture \cite{14}, and compression and shear \cite{15}. Topology based approach has been also considered, with focus on the contact network topology in isotropically compressed \cite{10} and tapped granular media \cite{17}.

We have recently started employing algebraic topological techniques for the purpose of quantifying forces in a manner which is global in character, but at the same time includes detailed information about the geometric structures of the forces. In \cite{18} we analyzed the number of components and loops, measured by Betti numbers (to be described in more detail below), as a function of compression (packing fraction) and the force level. That work considered the force networks on particle level: essentially, a total force on each particle was computed, and then the features of this force field were considered by analyzing the number of clusters (components), related in a broad sense to so-called force chains, as well as the number of loops, related to force chains’ connectivity. Realizing that a more complete description of a system can be reached by considering particle-particle interactions explicitly, we turned our attention to force networks whose basic building blocks are the interaction forces at particle contacts. Governed still by the idea of considering global properties, we have used persistent homology \cite{19}, which is best thought of a continuous map that provides for a substantial reduction of data while preserving geometric structures. The output of persistent homology consists of persistence diagrams, PD’s, that provide a consistent identification of the force thresholds at which geometric features made up of sets of interacting particles appear and disappear. Persistence analysis has shown explicitly and on a global level that (i) the geometry of the forces between particles has different properties from the geometry of the contact networks and (ii) the properties of these forces may depend strongly on the material properties, such as friction or polydispersity, for the specific case of granular particles \cite{20}.

Due to the complexities involved in the studies of force networks based on any of the approaches discussed so far, most of the existing results have concentrated on analysis of these networks for static systems, and there are very few attempts to analyze dynamical aspects of the force networks, or even of time-dependent properties of the forces experienced by particles, see e.g., \cite{21} and the references therein. Clearly, dynamical aspects are of significant importance, since many of the systems of interest are time dependent, and one would like to understand how the networks evolve in such a setting; some recent examples in the literature where the evolution of interaction networks is clearly of importance include wave propagation through particulate systems \cite{13} and impact \cite{22}.

There are many questions related to temporal evolution of force networks that one could ask, such as: What are the generic features of the temporal evolution of force networks? Are these features different for unjammed, but dense systems compared to the jammed ones? What happens as a system goes through jamming transition? What is the influence of friction on the evolution of force networks? We will address some of these questions in the present work. The tools that we will use involve the concept of ‘distance’ between PD’s, measuring (in a manner that will be made precise) the amount of change in force networks from one state to the next. To be specific, we will consider a particular system of inelastic frictional particles, but the technique that will be described is independent of the model describing particle-particle interaction, and could be equally well applied to the systems of particles of different shapes interacting by any other means.

The remainder of this paper is organized as follows. In Sec. \ref{sec:sys} we discuss the system to be studied. A brief introduction to persistence is given in Sec. \ref{sec:background} more in-depth analysis is given in \cite{23}. In this section we also introduce the concept of distance between the persistence diagrams. In Sec. \ref{sec:results} we present the results of persistence analysis. Section \ref{sec:discussion} summarizes the results and discusses possible future directions.

\section{The Systems to be Studied} \label{sec:sys}

We focus on the system of soft inelastic possibly frictional disks in two spatial dimensions (2D), bound by inward moving rough walls, similarly as in our earlier works \cite{18,20}. The simulations are of the type utilized in \cite{18} for completeness, a brief outline is given in Appendix \ref{app:method} and a short description follows.

The inward motion of wall particles is slow as described in the Appendix, so that the energy provided by the walls’ motion is dissipated quickly and therefore compression waves or other type of large scale inhomogeneities are not present in the system: the packing fraction, $\rho$, is essentially uniform, except possibly for lower $\rho$’s, which are not the focus of this study. The particles are characterized by the coefficient of Coulomb friction, $\mu$, and polydisperse, with a random distribution of diameters in the range $1 \pm r_p/2$ scaled by the average particle diameter, $d_{\text{ave}}$; we use $r_p = 0.4$. The initial size of the domain considered is $L^2$, with $L = 50 d_{\text{ave}}$. The initial configuration is formed by placing particles on the rectangular
lattice and then giving them initial random velocities. After this initially supplied energy is dissipated (through friction and inelasticity), the system is compressed leading to ρ’s in the range [0.63 : 0.90]. Gravitational effects are not considered.

As the system is compressed, we extract the current values of the forces between the particles at specified time intervals. To provide visual picture of the system that is being analyzed, Fig. 1(a-c) shows the magnitude of the normal forces acting between the particles for three nearby values of ρ (animations of the evolution of the force network as the system is compressed are available as Supplementary Materials of [20]). The forces are normalized by the average normal force at the given time. Note that the figures appear very similar, and additional inspection shows that both the average contact number, Z, and the average force are almost identical. Differences between the networks become visible if we plot changes in the magnitude of the normal force. Figure 1(d-e) shows these differences, and we can see that the difference between (a-b) is localized, while the (a-c) differences have more of a global character. The main focus of this paper is to introduce and discuss a set of measures that can be used to quantify these differences in a precise manner.

III. METHODS

We start by describing force networks in the manner that will allow us to use topological techniques for their quantification. As we will see, important features of the force network can be captured by using persistence diagrams, PD’s, that were already introduced in our previous work [20], and discussed in depth in [23]. A brief overview of persistence is given in Sec. III A followed by a couple of examples in Sec. III B and discussion of PD’s for the simulation data in Sec. III C. Then, in Sec. III D we discuss different types of metric on the space of PD’s. The main feature of these metrics is different sensitivity to various changes of the force network. This makes them useful for distinguishing local and global changes, as we demonstrate by analyzing the networks shown in Fig. 1.

The computations that are outlined in this Section and used for the rest of the paper are based on publicly available open source software [24, 25].

A. Force Networks

We encourage the reader to view the images shown in Fig. 1 as 2D landscapes with the altitude given by the normal force magnitude at a particular spatial point. The goal of this section is to briefly recall the basic elements of persistent homology which we use to provide a concise characterization of the geometry of these landscapes.

We begin by making precise the process by which we obtain Fig. 1(a - c). Given a collection of particles \{pi | i = 0, ..., N\}, we define a simplicial complex CNi called the interaction complex consisting of vertices \{vi | i = 0, ..., N\} where each vertex vi is identified with particle pi and all possible edges \{vi, vj\}. Now let ψi,j ∈ ℝ denote the magnitude of the force experienced between particles pi and pj, then the function f is defined on the edges by

\[ f(⟨vi, vj⟩) := ψi,j. \]

This function is extended to the vertices by

\[ f(vi) = \max_{j=0, ..., N} \{f(⟨vi, vj⟩)\}. \]

Figure 1(a - c) indicates the value of \( f(v_i, v_j) \) for which \( f(⟨v_i, v_j⟩) > 0 \).

Observe that in these figures there are triangles made up of three edges. These represent loops in the contact network made up of three particles. Since in a perfect densely packed crystalline structure made up of disks of the same size all loops would be made up of exactly 3 particles, we refer to a loop involving four or more particles as a defect. In the analysis we perform in this paper we have chosen to focus on defects. To avoid counting the three particle loops we extend CNi to its flag complex CNi by adding all triangles \{v_i, v_j, v_k\} and defining

\[ f(⟨v_i, v_j, v_k⟩) = \min \{f(⟨v_i, v_j⟩), f(⟨v_i, v_k⟩), f(⟨v_j, v_k⟩)\}. \]

Since our goal is to consider networks of contacts with forces larger than a given force, we are interested in the geometry of a part of the complex on which the forces exceed a specified level. In particular, given a threshold θ > 0, define the associated interaction force network

\[ \text{FN}_i(f, θ) := \{σ ∈ \text{CN}_i | f(σ) ≥ θ\}, \]

which corresponds to the part of the contact network experiencing force larger than θ. Since we are working with finite sets of particles, the function \( f \) can take on a finite set of values \( Θ = \{θ_n\} \). The interaction force network filtration is the collection of interaction force networks \( \{\text{FN}_i(f, θ) | θ ∈ Θ\} \). For simplicity we drop the adjective interaction and write force network and force network filtration.

B. Persistent Homology: an example

In the context of the two dimensional simulations that are studied in this paper the simplest geometric structures that can be quantified by persistent homology are connected components and loops. We use the simple two-dimensional example shown in Fig. 2 to provide some intuition to what these measurements represent. The reader is referred to [23] for a more complete description, along with a simpler one-dimensional example. Figure 2 shows two threshold values, 86 (a) and 55 (b). The function (along with the associated threshold level) for which persistent homology is being computed is shown.
FIG. 1: (a-c) Snapshots of the simulated system \((r_p = 0.4 \mu = 0.5)\) at three values of \(\rho\) close to 0.79. Every edge in the network corresponds to the normal force acting between the particles and the color encodes the magnitude of the normal force normalized by the current value of the average normal force. (d-e) The absolute values of the differences between the normal forces shown in (a-b) and (a-c), respectively (note different color scheme compared to (a-c)).

in the upper left hand corner of (a) and (b). The upper right hand corners show the associated the force networks, \(FN_I(f, 85)\) and \(FN_I(f, 55)\). Each panel contains a \(\beta_0\) PD, denoted by \(PD_0\), which measures connected components, and a \(\beta_1\) PD, denoted by \(PD_1\), which measures loops. Note that the PD’s in (a, b) are the same.

Since we are using super level sets to define the force networks, we compute the PD’s by descending through the threshold levels. For the moment let us focus on Fig. 2(a) associated with the threshold \(\theta = 85\). The force network \(FN_I(f, 85)\) contains no loops. This can be seen in \(PD_1\) by noting that there are no points in the diagram to the lower right of (85, 85). There are, however, four components and in the \(\beta_0\) diagram to the lower right of (85, 85) there are four points: (128, -1), (119, 74), (113, 83), and (106, 76). We remark (this information is not contained in the \(PD_1\)) that the point (128, -1) corresponds to the lower component, (119, 74) to the upper right component, (113, 83) to the upper left component, and (106, 76) to the middle component. These points contain important information: the first (birth) coordinate indicates the threshold at which the component appears and second (death) coordinate indicates where the component merges with another older component. In particular, we can now conclude that the first point in the lower component appeared at threshold level 128, the first point in the upper right component at threshold level 119, etc.

To explain the comment about the ”older component” note that first component to appear at threshold level 128. Since this is the first component it is always the oldest and thus never disappears. This is indicated by assigning a death coordinate of -1. The death coordinate of the upper left component is 83 which implies
that at threshold 83 it merges with another component. The geometry suggests that it does not merge with the lower component at that threshold. Thus, it can merge with the upper right component or the middle component. However, the birth value of the middle component is 106 which makes it younger than the upper left component. The upper right component has birth value 119 which makes it older. Thus when the upper right component merges with the upper left component at threshold 83, the upper left component dies.

Turning to Fig. 2(b) associated with threshold $\theta = 55$, we note that to the lower right of $(55, 55)$ there are two points in the $\PD_0$ and three points in the $\PD_1$, indicating that the associated force network $\FN_I(f, 55)$ has two components and three loops, respectively. Observe that the lower loop has just formed and thus the associated birth coordinate is 55. The death value occurs at the threshold where a loop is filled in.

The convention that the younger feature dies has an important implication. Let $(\theta_b, \theta_d)$ be a point in a PD. The number $\theta_b - \theta_d$ is called the lifespan of the geometric feature associated with $(\theta_b, \theta_d)$. Observe that features with a longer lifespan persist over a longer range of values and hence are more robust. Conversely features with very short lifespans are often regarded as noise, since they persist only over a small range of force values and thus can be introduced by small perturbations.

Viewing the complete animation of Fig. 2 provided in the Supplementary Materials should convince the reader that persistent homology provides a concise encoding of the dominant geometric features of a function. The fact that it is concise implies that information is lost and thus two distinct functions can have the same persistence diagram. Of course, if two functions have distinct persistence diagrams, then they must exhibit distinct geometric features.

C. Persistence diagrams for simulation data

Having discussed the simple example of the previous section, we are now equipped to consider PD’s resulting from DES. Figure 3 shows the $\PD_0$ and $\PD_1$ diagrams for the force network defined on the flag complexes corresponding to Figs. 1(a) (the diagrams corresponding to Figs. 1(b - c) visually appear very similar and are not shown since they do not provide any additional information). Not surprisingly, these are more complex than those of Fig. 2. We begin our analysis with some simple observations (see [20] and [23] for a more detailed discussion).

- The $\PD_0$ show a ‘cloud’ of points in the $[0 : 3]$ birth range, meaning that most of the features (force chains, loosely speaking) start appearing at the force level which is about 3 times the average force, and disappear by $[0.8 : 1]$, suggesting that at a force level slightly smaller than the average force, most features disappear (merge);
- Careful inspection of $\PD_0$ shows two points with a death value of $-1$ level; one born at high force threshold $\theta = 3$, and the other at the zeroth force level. The interpretation of the former one follows from the convention discussed in the previous section that the component that is born first dies last; the latter is due to the presence of isolated particles (rattlers) or particles that experience only contacts with zero force. Rattlers can be detected because only the higher dimensional simplicies for which
the function \( f \) is positive are used in the persistent homology computations. Hence the rattlers create separate connected components at the zero force level.

- Note that for \( \text{PD}_1 \)'s a death level of \( \theta_d = -1 \) implies the existence of a defect. If \( \theta_d > 0 \), then the hole is filled in, and \( \theta_d \) indicates the weakest magnitude of interacting forces within the region enclosed by the loop. The \( \text{PD}_1 \) in Fig. 3(b) suggest that the birth value \( \theta_b \) of most of the loops are less than about 1.5 the average force. That the birth values of the \( \text{PD}_1 \) is lower than that of the \( \text{PD}_0 \) is not surprising; the birth of a loop corresponds to the lowest magnitude of the normal forces acting along the contacts forming the loop. It is also worth noting that most of the loops are associated with defects.

While these observations about individual PD's are of interest, it is clear that a more systematic approach is needed to quantify and interpret the available information. This is especially true in the context of comparing time dependent systems, where each time step can produce a distinct PD. To make this comparison precise requires the introduction of appropriate metrics to measure the difference between PD's.

**D. Distance between persistence diagrams**

At a minimum, an appropriate metric for PD’s must satisfy the property that if two functions are similar, then the distance between the associated PD’s must be small. It was shown in [19] that such metrics exist. Explicit formulas and a detailed discussion in the context of DGM is presented in [23, Definition 7.1]. However, since the precise definitions are somewhat technical, we limit ourselves to a heuristic presentation. Consider Fig. 4(a) where one can consider the function \( g \) as a noisy perturbation of \( f \). To understand the metrics we recall two observations from Sections III.B and III.C: (1) if the points in two PD’s lie in the similar regions then the PD’s should be close, and (2) short lifespans, i.e. persistence points near the diagonal, are related to small perturbations. This suggests that given two PD’s one attempts to match points from one diagram with points in the other diagram or points on the diagonal in such a way as to minimize distances between the matched points. Figure 4(a) suggests such a minimizing matching. Let \( \gamma : \text{PD}_n(f) \rightarrow \text{PD}_n(g) \), denote such a matching between two PD’s. The *bottleneck distance* is denoted by \( d_B(\text{PD}_n(f), \text{PD}_n(g)) \) and is defined by \( \sup_{p \in \text{PD}_n(f)} \| p - \gamma(p) \|_\infty \), while the degree-q Wasserstein distance is denoted by \( d_{W^q}(\text{PD}_n(f), \text{PD}_n(g)) \) and is defined by \( \left( \sum_{p \in \text{PD}_n(f)} \| p - \gamma(p) \|_\infty^q \right)^{1/q} \) where in both cases \( \gamma \) is chosen to minimize these quantities.

In this paper we restrict ourselves to the bottleneck distance \( d_B \) and the Wasserstein distances \( d_{W^q}, q = 1, 2 \). Observe that the bottleneck distance reports only the single largest difference between PD’s while \( d_{W^q} \) includes all differences between the diagrams. Thus, it is always true that

\[
d_B \leq d_{W^q}.
\]

Sensitivity of the Wasserstein distances to small differences (possibly due to noise) can be modulated by the choice of the value of \( q \), i.e. \( d_{W^2} \) is less sensitive to small changes than \( d_{W^1} \).

Example shown in Fig. 3(a) demonstrates the differences between \( d_B \) and \( d_{W^1} \). Let \( \text{PD}_0(f) \) and \( \text{PD}_0(g) \) be the \( \text{PD}_0 \)'s corresponding to the functions \( f \) and \( g \), respectively. Fig. 3(b) shows that the \( \text{PD}_0(f) \) consist of a single point \((5, -1)\). The \( \text{PD}_0(g) \) is more complicated. There is one copy of \((3, -1)\) corresponding to the dominant feature of \( g \) and seven copies of \((1, 0)\) representing seven smaller features. Since the bottleneck distance considers only the largest feature, \( d_B(\text{PD}_0(f), \text{PD}_0(g)) = 2 \). On the other hand, \( d_{W^1}(\text{PD}_0(f), \text{PD}_0(g)) = 5.5 \) since the contribution of every small feature to the \( d_{W^1} \) distance is 0.5 and \( d_{W^2}(\text{PD}_0(f), \text{PD}_0(g)) \approx 2.6 \). Thus the different metrics distances provide complementary information about the differences between two functions (or two landscapes). In particular, the fact that the \( d_{W^2} \) distance is closer to \( d_B \) than \( d_{W^1} \) suggests that the geometric differ-
than the relative difference between distances are larger than those of the bottleneck. What matters here is that the relative difference between the geometries between these landscapes. Note that this does not imply that there are no significant point wise differences (Fig. 1(d-e)), that measures the point wise difference, extends to 0.4 (40% of the average force). Combining these two observations suggests that the point wise locations of the strong and weak particle interactions have shifted, but their relative geometries have remained similar.

As is to be expected the values of the Wasserstein distances are larger than those of the bottleneck. What is worth noting is that the relative difference between $d_{W1}(P^a, P^b)$ and $d_{W1}(P^c, P^d)$ is significantly larger than the relative difference between $d_{W2}(P^a, P^b)$ and $d_{W2}(P^c, P^d)$. This suggests that the force landscapes of Fig. 1(c) differs from the one of Fig. 1(a) via many more small geometric changes than the force landscape of Fig. 1(b) differs from that of Fig. 1(a). This is consistent with the difference plots of Fig. 1(d-e) which show that in general the values of Fig. 1(e) are higher than those of Fig. 1(d) and spread over a broader range of the domain.

TABLE I: Distances between the persistence diagrams for the force networks corresponding to Fig. 1(a-c).

|     | $\beta_0$ | $\beta_1$ |
|-----|------------|------------|
| $(P^a, P^c)$ | $d_B$ | $d_{W1}$ | $d_{W2}$ |
| $d_B$ | 0.089 | 0.081 | 3.8 | 0.57 |
| $W_{d_B}$ | 0.12 | 14.0 | 0.89 | 0.15 | 19.2 | 0.99 |

TABLE II: Distances between the persistence diagrams for the different parts of the force networks corresponding to Fig. 1(a,b).

|     | $\beta_0$ | $\beta_1$ |
|-----|------------|------------|
| $(P^a, P^b)$ | $d_B$ | $d_{W1}$ | $d_{W2}$ |
| $d_B$ | 0.16 | 0.13 | 5.8 | 0.59 |
| $W_{d_B}$ | 0.18 | 0.14 | 5.6 | 0.50 |
| $d_B$ | 0.10 | 0.12 | 5.5 | 0.51 |
| $W_{d_B}$ | 0.11 | 0.087 | 5.4 | 0.87 |

TABLE III: Distances between the persistence diagram for the different parts of the force networks corresponding to Fig. 1(a,c).

TABLE IV: Result ...

So far, we have shown that the various distances can be used not only to quantify the differences between force networks, but also to distinguish between local and global differences; we will use these findings to analyze the results of DES in the rest of this paper.

Before concluding this section, we illustrate that the distances can be also used to isolate a part of the domain where dominant differences are present, although we will not use this approach in the rest of the paper.

Consider again the networks shown Fig. 1(a-c). Divide each domain into four equal blocks: bottom left, bottom right, top right and top left. Table III gives the distances between corresponding blocks for the (a-b) PD’s. We see that the distances between the bottom left blocks are always larger than the distances between the other three blocks. Hence we can conclude that differences between the networks FN$_i^b$ and FN$_j^b$ are concentrated in the bottom left corner. Table III shows the distances between the four blocks for the networks FN$_i^b$ and FN$_j^b$. Here, the values are very similar for all the blocks. This finding indicates that the changes are distributed evenly over the entire domain.

So far, we have applied the distance concept to discuss the differences between small number of force networks. Now we will proceed to analyze large number of force networks describing states of the systems as they are evolve through the jamming transition.

IV. RESULTS

In the DES that we have carried out, we slowly compress the system through a range of packing fractions $\rho = [0.63 : 0.90]$ as discussed in Sec. I. While the system is being compressed, we output the force information at approximately fixed time intervals. In the present work...
we focus on results obtained by extracting approximately 100 samples, and averaged over 20 realizations for the purpose of obtaining statistically significant results.

We compute the distance between PD’s computed using the force information from consecutive samples. Since the PD’s provide information about the geometry of the forces, it is useful to consider the plots presented in this section as a measure of the rate of change of the geometry of the forces.

In what follows, we focus first on the normal forces in a polydisperse frictional system \((r_p = 0.4, \mu = 0.5)\) and then proceed to discuss the tangential forces in this system, as well as a polydisperse frictionless system \((r_p = 0.4, \mu = 0)\).

### A. A polydisperse frictional system

Figure 6 shows various distances between the PD’s for consecutive (normal) force networks, averaged over 20 realizations for frictional polydisperse system \((r_p = 0.4, \mu = 0.5)\). We also plot the average number of contacts per particle, \(Z\). The most pronounced feature is the dramatic change around the jamming point, \(\rho_J\), loosely defined as the \(\rho\) at which \(Z \approx 3\) \([18]\). Each distance, however, behaves differently as the system is compressed; as demonstrated below, these differences provide us with additional insight regarding the evolution of forces as the system goes through the jamming transition.

Before comparing individual distances let us put their values into a context. Recall that the normal force used for computing PD’s is normalized by the current average value. This average is small for \(\rho < \rho_J\) and increases (approximately linearly) after the jamming transition. Using the landscape analogy there are at least two ways in which to interpret \(d_B = 1\). The first is that a geometric feature with lifespan equal to average force has either appeared or disappeared. The second is that the length of the lifespan of a single geometric feature has changed by the average force.

We begin our analysis by considering the evolution of the forces prior to jamming, \(\rho < \rho_J\). Figure 6(a) gives the distances between consecutive PD’s, therefore showing the rate of evolution of connected components (force chains in the loose sense). The reported rate clearly depends on the metric being used. The relatively large values \(d_B\) is an order of magnitude greater than one for most of this regime) suggest rapid changes in the structure of the forces. There are at least two types of events that are monitored; since there is still considerable room for particles to move, we are observing the collisions of particles that create large spikes in the forces, and we are observing changes in the connectivity of the system as particles come together, create a larger connected component with relatively strong force interactions inside this component, and then separate again. The \(d_B\) metric is capturing the single largest of these events while \(d_{Wq}\), \(q = 1, 2\), are measuring changes of the forces globally.

Figure 6 plots the relative rates of change \(d_B/d_{W1}\) and \(d_B/d_{W2}\). These relative rates of change are roughly constant for \(\rho < 0.75\). Furthermore, \(d_B/d_{W2} \approx 0.75\) and \(d_B/d_{W1} \approx 0.2\) suggesting that from frame to frame there is a single large event and a number of smaller events that contribute to the change in geometries.

At \(\rho \approx 0.75\), the behavior of the system begins to change. The rate of change as measured by \(d_B\) decreases dramatically, suggesting absence of collisions characterized by large relative velocity of the particles, or of large changes in connectivity of the system accompanied by an increase of the forces acting between the colliding components. For \(0.75 < \rho < \rho_c\) the ratio of \(d_B/d_{W2}\) continues to be roughly constant until just before the jamming when it plunges. This suggests that there continues to be a single large event. However, over this same
range the rate of change as measured by $d_W^1$ dramatically increases (Fig. 6(a)) and hence the ratio of $d_B/d_W^1$ decreases, suggesting that the number of small events is growing. The physical interpretation is that as one approaches the jamming transition, there is a rapid reorganization of the structure of the forces, taking place globally through many small rearrangements.

Visual inspection of Fig. 6(a-b) shows that there is a significant difference between the rate of change in the geometries as measured by PD$_0$ and PD$_1$. For $\rho < \rho_J$, this is not surprising. The particles have room to move around. Thus, the number of loops is small and furthermore, since it is unlikely that all the forces between the particles forming loops are strong, the points in the PD$_1$ tend to appear for smaller forces and have a shorter life span.

With this in mind, it is perhaps not surprising that before jamming the rate of change of the loop structure generally increases as a function of $\rho$. At $\rho \approx 0.77$ the $d_B$ distance starts decreasing. We interpret this to mean that if the system is sufficiently packed then it becomes difficult to support the appearance or disappearance of loops with large lifespans or for which the maximal or minimal forces change dramatically. It should be noted, however, that while the size of the individual changes becomes constrained, the locations at which changes can occur do not. As it can be seen from Fig. 7, both ratios, $d_B/d_W^2$ and $d_B/d_W^1$, begin to decrease at $\rho \approx 0.76$. The fact that $d_B/d_W^2$ decreases suggests that multiple large changes in the loop structures are occurring, along with the increase in the number of small changes, as indicated by $d_B/d_W^1$. We conclude therefore that during the jamming transition there is a significant global reorganization of the loop structure with a variety of local changes of the magnitude comparable to the largest one.

The analysis past the jamming transition, $\rho > \rho_J$, is simpler; the rate of change of the forces of the system slows dramatically. This is not surprising. A large part of the contact network is fixed, thus the particles cannot make and brake contacts, nor can the magnitudes of the normal forces change dramatically, thus, for example, the rate of change measured by $d_B$ must be small. What is more interesting to note is that the ratios of $d_B/d_W^2$ and $d_B/d_W^1$ remain essentially constant throughout this regime and are roughly one and three orders of magnitude smaller, respectively, than before the jamming transition. This implies that even though the large events as measured by $d_B$ are getting smaller, they are also becoming more broadly distributed ($d_B/d_W^2$ smaller by one order of magnitude) and there are many more small events ($d_B/d_W^1$ smaller by three orders of magnitude).

### B. Tangential forces

While in the research related to force networks the focus is usually on normal forces, it is appropriate to ask whether evolution of tangential forces provides any additional information. Figure 8 shows the corresponding results, where now the forces are normalized with the average tangential force (as expected, the tangential force average is significantly smaller than the normal one). Perhaps the most interesting feature of the results shown in Fig. 8 is how similar they are to the results obtained for normal forces, shown in Fig. 6(except perhaps very close to $\rho_J$). This finding suggests that both for $\rho < \rho_J$ and for $\rho > \rho_J$, the evolution of tangential forces to large degree follows the evolution of normal forces, both regarding the connected components and the loops. For $\rho \approx \rho_J$, we see larger deviations between normal and tangential forces, with a particular feature that the distances that include all the differences between the networks, such as $d_W^1$ and $d_W^2$, are significantly smaller for tangential forces,
suggestions less dramatic evolution of these forces as the system goes through jamming transition. The bottleneck distances, $d_B$, show however similar features and magnitudes for the normal and tangential forces throughout the evolution, including $\rho \approx \rho_J$, suggesting that the largest change in the tangential forces is slaved to the largest change in the normal ones.

C. A polydisperse frictionless system

Finally, we discuss briefly the normal force network for polydisperse frictionless ($\mu = 0$) system. In earlier work [20], we discussed the information that can be obtained by considering the PD’s, and found, based on the number of generators, that the force networks for $\mu = 0$ systems appeared to be extreme, in the sense that the number of generators for frictionless system was significantly larger (see Fig. 6 in [20]), compared to frictional systems. By considering the distances between the states of the systems, we can now discuss how friction influences the evolution of force networks.

Figure 8 shows the corresponding distances and the $Z$ curve. Note that the jamming transition is shifted to larger $\rho$’s for $\mu = 0$ systems [20]. Considering connected components, Fig. 9(a), we make two observations.

- For $\rho < 0.65$ the rate of change for $\mu = 0$ systems, as measured by $d_B$ and $d_W$, is roughly the same as for $\mu \neq 0$ systems. However, $d_W$ is approximately 40% smaller for $\mu = 0$. In fact, as is seen by comparing Fig. 10(a) and Fig. 7(a) over the entire range of $\rho$ adjusted for $\rho_J$, the ratio $d_B/d_W$ is consistently larger for the $\mu = 0$ system while $d_B/d_W$ is roughly the same. This suggests that friction plays a role in the appearance of small events.
- Initially, as $\rho$ increases, the rates of change increases more rapidly in the frictional system. However, both $d_B$ and $d_W$ begin slowing down at a much lower $\rho$ relative to
\[ β_0, \frac{dB}{dW_1}, \frac{dB}{dW_2}, \beta_1, \frac{dB}{dW_1}, \frac{dB}{dW_2} \]

\( (a) \beta_0, \quad (b) \beta_1 \)

FIG. 10: The ratios between the rates of change as measured by different metrics for polydisperse frictional systems (\( r_p = 0.4, \mu = 0 \)).

\( ρ_J \) in the frictional system, suggesting that as the particles come in closer contact, friction limits the magnitude of the largest changes in the geometry.

As indicated above, there are more points in the PD1 for \( \mu = 0 \) than for \( \mu = 0.5 \). Thus the fact that for \( ρ \) significantly less than \( ρ_J \), the rate of change is much less as measured by \( dB \) and \( dW_2 \) in the \( \mu = 0 \) system as opposed to the \( \mu = 0.5 \) system (compare Fig. 9(b) and Fig. 6(b)), strongly suggests that friction plays an important role in creation, destruction, and deformation of the loops when the system is compressed but not yet jammed. Observe that close to jamming, the rate at which the loop structure changes for the \( \mu = 0 \) systems is significantly more dramatic, both regarding the largest change, measured by \( dB \), and regarding overall changes, measured by \( dW_1 \) and \( dW_2 \). Even after jamming, for \( ρ > ρ_J \), the rate of change of the \( \mu = 0 \) systems is approximately twice that of the \( \mu = 0.5 \) systems. It is interesting that the influence of friction for \( ρ > ρ_J \) is much more obvious for loops than for connected components. Further research is needed to better understand this finding.

V. CONCLUSIONS

The evolution of force networks in dense particulate systems presents a complicated problem that is difficult to analyze using conventional techniques. In this paper we show that computational homology can be used to address many aspects of the problem. In particular it provides a unique set of precise and well defined measures of the evolution of the geometry of the inter particle forces. The main findings that apply to frictional and frictionless systems, to normal and tangential forces, to both connected components (force chains) and loops, and to all different distances between the considered states, are as follows:

- The comparison of considered measures describing the evolution of force networks shows that for unjammed systems, large changes of the considered force networks are possible, but these changes typically consist of local, isolated events. However, as the system goes through the jamming transition, the measures that we have implemented suggest large changes of the force networks on global, system-wide scale;
- The evolution of force networks is considerably different between unjammed, but dense systems, and the jammed ones. For jammed systems, the evolution of the networks is significantly less dramatic, in the sense that the distances between the consecutive states of the system are significantly smaller.

In addition to these general findings, we also list the main findings that focus on the specific forces/systems/distances:

- The evolution of tangential forces is similar in its main features to the evolution of the normal ones, with some differences particularly close to \( ρ_J \), where jamming occurs;
- There are significant differences in the evolution of force networks for frictionless versus frictional systems. With regard to the component structure of the force network (recall that this is related in a broad sense to so-called force chains) the peak rates of evolution before jamming are 20 – 40% higher for the frictional system than for the frictionless system, but become quite similar after jamming. The differences are even more pronounced when the evolution of loop structures is considered. For \( ρ \ll ρ_J \), the rate of evolution measured by \( dB \) and \( dW_2 \) is larger for the frictional system. For \( ρ \lesssim ρ_J \), the relative rates change and thereafter the difference between the considered states are significantly more pronounced in frictionless systems.

In this work, we have concentrated on describing the
concept of distance between the persistence diagrams in the context of force networks, and have analyzed a particular set of systems (2D polydisperse circular particles with/without frictional effects) exposed to a slow compression and sampled with prescribed sample rate. Furthermore, we have focussed on the generic, averaged features of the differences between considered force networks. In future work, we will consider in more detail what is the influence of particle shape, sampling rate, and of the protocol used to evolve the considered particulate system, as well as analyze more precisely the evolution of networks in single realizations [27].

Before closing, we note that one particular strength of the computational homology approach is that it is system-independent and can be applied to any particulate system, independent of the physical properties of the considered system, such as the type of interaction between the particles, or dimensions of the physical space in which they live. Therefore, the natural next step is to consider particulate systems characterized by different physical properties (cohesion, shape) or different geometry (2D versus 3D). Furthermore, the approach can be as well applied to experimental systems, such as those built from photoelastic particles, where detailed information about the force networks is available. Analyses of such systems will be the subject of our future works.

Acknowledgments: We thank Vidit Nanda for valuable technical assistance with Perseus. We acknowledge the support by NSF DMS-0835611, DTRA 1-10-1-0021 (LK); and NSF DMS-0915019, CBI-0835621 and contracts from DARPA and AFOSR (AG, MK, KM).

Appendix A: Discrete Element Simulations (DES)

In the simulations, circular grains are confined to a square domain with rough walls composed of monodisperse particles. The walls move inward at constant speed, \( v_c \). No annealing of the system is carried out, and gravity is neglected. The particle-particle (and particle-wall) interactions include normal and tangential components. The normal force between particles \( i \) and \( j \) is

\[
F^n_{i,j} = k_n n_i n_j - \gamma_n \bar{m} v^n_{i,j} ,
\]

where \( r_{i,j} = |r_{i,j}| \), \( r_{i,j} = r_i - r_j \), \( n = r_{i,j}/|r_{i,j}| \), and \( v^n_{i,j} \) is the relative normal velocity. The amount of compression is \( x = d_{i,j} - r_{i,j} \), where \( d_{i,j} = (d_i + d_j)/2 \), \( d_i \) and \( d_j \) are the diameters of the particles \( i \) and \( j \). All quantities are expressed using the average particle diameter, \( d_{ave} \), as the lengthscale, the binary particle collision time \( \tau_c = \pi \sqrt{d_{ave}/(2gk_n)} \) as the time scale, and the average particle mass, \( \bar{m} \), as the mass scale. \( \bar{m} \) is the reduced mass, \( k_n \) (in units of \( mg/d_{ave} \)) is the spring constant set to a value that corresponds to that for photoelastic disks [28], and \( \gamma_n \) is the damping coefficient [29]. The parameters entering the linear force model can be connected to physical properties (Young modulus, Poisson ratio) as described e.g. in [29].

We implement the commonly used Cundall-Strack model for static friction [30], where a tangential spring is introduced between particles for each new contact that forms at time \( t = t_0 \). Due to the relative motion of the particles, the spring length, \( \xi \) evolves as

\[
\xi = \int_{t_0}^{t} v^t_{i,j} (t') \, dt' ,
\]

where \( v^t_{i,j} = v_{i,j} - v^n_{i,j} \). For long lasting contacts, \( \xi \) may not remain parallel to the current tangential direction defined by \( t = v^t_{i,j}/|v^t_{i,j}| \) (see, e.g., [31]); we therefore define the corrected

\[
\xi' = \xi - n (n \cdot \xi) ,
\]

and introduce the test force

\[
F^t = \min(\mu_s |F^n|, |F^t'|) F^t'/|F^t'| ,
\]

and redefine \( \xi \) if appropriate.

For the initial configuration, particles are placed on a square lattice and given random initial velocities; we have verified that the results are independent of the distribution and magnitude of these initial velocities. The wall particles move at a uniform (small) inward velocity \( v_c = 2.5 \cdot 10^{-5} \). We integrate Newton’s equations of motion for both the translation and rotational degrees of freedom using a 4th order predictor-corrector method with time step \( \Delta t = 1/50 \). In this work, we consider systems with \( N = 2000 \) particles with \( k_n = 4 \cdot 10^3 \), \( e_n = 0.5 \), \( \mu_s = 0.5 \), and \( k_l = 0.8k_n \).

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