The arrangements of particles and forces in granular materials have a complex organization on multiple spatial scales that range from local structures to mesoscale and system-wide ones. This multiscale organization can affect how a material responds or reconfigures when exposed to external perturbations or loading. The theoretical study of particle-level, force-chain, domain and bulk properties requires the development and application of appropriate physical, mathematical, statistical and computational frameworks. Traditionally, granular materials have been investigated using particulate or continuum models, each of which tends to be implicitly agnostic to multiscale organization. Recently, tools from network science have emerged as powerful approaches for probing and characterizing heterogeneous architectures across different scales in complex systems, and a diverse set of methods have yielded fascinating insights into granular materials. In this article, we review work on network-based approaches to studying granular matter and explore the potential of such frameworks to provide a useful description of these systems and to enhance understanding of their underlying physics. We also outline a few open questions and highlight particularly promising future directions in the analysis and design of granular matter and other kinds of material networks.

Keywords: granular materials; particulate systems; networks; multiscale organization.
Glossary of Terms

Granular materials: Granular materials are collections of discrete, macroscopic particles that interact with each other through contact (rather than long-range) forces. Importantly, these systems are non-equilibrium: the particles are large enough to avoid rearrangement under thermal fluctuations, and they lose energy through frictional and inelastic interactions with neighboring particles.

Particulate materials: Like granular materials, particulate materials are collections of discrete, macroscopic elements. However, the elements making up the system may be entities—such as bubbles, foams, colloids, or suspensions—that include multiple phases of matter. The term “particulate material” is a more general one than “granular material”.

Packing fraction: The fraction of a granular material that is physically occupied by the particles. One calculates the packing fraction as the ratio of the total volume of all particles to the total volume of the region that contains the particles. The packing fraction is also sometimes called the “packing density” or “volume fraction”.

Force chain: Force chains are typically described as the subset of inter-particle contacts in a granular material that carry the largest forces in the system. They often form filamentary networks that align preferentially with the principal stress axes under which a material is loaded.

Jamming: As certain system parameters change, disordered, particulate materials typically undergo a transition from an underconstrained, liquid-like state to a rigid, solid-like state characterized by the onset of mechanical stability. The transition to/from a jammed state often arises through increases/decreases in packing fraction (or in contact number), which can occur due to an applied load. A formal theory of jamming exists for idealized situations (with soft, frictionless, and spherical particles).

Isostatic: A jammed packing is isostatic when it has exactly the minimum number of contacts that are required for maintaining mechanical stability through force balance and torque balance. One typically examines isostaticity by calculating the mean number of contacts per particle. “Hyperstatic” and “hypostatic” packings have more and fewer contacts, respectively.

Mono/bi/poly-disperse: A particulate material is monodisperse if it is composed of particles of a single species (i.e., particles with the same size, shape, and material properties). Bidisperse materials have particles of two species, and polydisperse materials can either have particles from three or more discrete species or have particles with a continuum of properties.

Structural rigidity theory: For a specified mechanical structure that is composed of fixed-length rods connected to one another by flexible hinges, structural rigidity theory studies the conditions under which the associated structural graph (in which the edges correspond to the rods and the nodes correspond to the hinges) is able to resist deformations and support applied loads.

Stress: A stress is a force applied to an object’s surfaces. (The units measure a force per unit area.) Shear stress arises from the component of the applied force that acts in a direction parallel to the object’s cross-section, and normal stress arises from the perpendicular component.

Strain: Strain is the fractional (unitless) deformation of a material that arises due to an applied stress. One calculates strain from the relative displacement of particles in a material, excluding rigid-body motion such as translation or rotation. Like stress, strain has both shear and normal components.

Pure shear: One can use the term “pure shear” to describe either stresses or strains in which an object is elongated along one axis and shortened in the perpendicular direction without inducing a net rotation.

Axial compression: In axial compression, one applies inward forces to an object in one direction (uniaxial), two directions (biaxial), or all directions (isotropic compression). These forces result in uniaxial strain, biaxial strain, or isotropic strain, respectively.
Cyclic shear/compression: These terms refer to procedures in which repeated cycles of shear or compression are applied to the same system.

Shear band: A shear band is a narrow region of a particulate material in which most of the strain is localized, whereas other regions remain largely undeformed. A shear band is also sometimes called a region of “strain localization”.

Strain softening/hardening: As a material is loaded and undergoes deformation, continuing deformation can become either easier (strain softening) or harder (strain hardening). Eventually, after much deformation, the material can reach a critical state in which there are no further changes in the resistance to deformations.

Stress ratio: The stress ratio, which is analogous to Coulomb’s Law, is the ratio of shear to normal stresses. Frictional failure occurs when the shear force exceeds the product of the normal force and the coefficient of friction.

Photoelasticity/birefringence: Photoelasticity is an optical technique for quantifying internal stresses in a material. This experimental methodology is based on the transmission of polarized light through “birefringent” materials, which have preferentially fast and slow directions for the propagation of light.

DEM or MD simulations: The Discrete (or Distinct) Element Method and Molecular Dynamics simulations are related numerical techniques that compute the motions of all particles in a system (such as in a granular material). In both methods, a computer algorithm treats each particle as an object subject to Newton’s laws of motion, where forces consist of body forces (e.g., gravity) and those that arise from interactions with the object’s neighbors.

1. Introduction

Granular materials comprise a subset of the larger set of particulate matter [1–6]. People engage with such materials—which include sands, beans, grains, powders such as cornstarch, and more—often in their daily lives. One can define a granular material as a large collection of discrete, macroscopic particles that interact only when in contact. Granular materials are inherently non-equilibrium in two distinct ways, characterized by (1) the lack of rearrangement under thermal fluctuations and (2) the loss of energy through frictional and inelastic dissipation during contact between grains. Nonetheless, they phenomenologically reproduce equilibrium states of matter, exhibiting characteristics of solids (forming rigid materials), liquids (flowing out of a container) or gases (infrequent contacts between grains), depending on the type and amount of driving. In this review, we focus mainly on granular solids and slow (non-inertial) flows [7]; these are dense materials in which sustained inter-particle contacts provide the dominant contribution to material properties. The functional properties of granular materials are related in a non-trivial way to the complex manner in which particles interact with one another and to the spatial scales (particle, chain, domain and bulk) and time scales over which those interactions occur. For example, pairs of particles can exert force on one another in a local neighbourhood. However, as particles push on adjacent particles, the combined effect can transmit forces over long distances via important mesoscale structures commonly called force chains [8, 9]. The idea of networks has been invoked for many years to help provide a quantitative understanding and explanation of force-chain organization [10–14]. Broadly speaking, force chains form a network of filamentary-like structures that are visually apparent in images from experiments, like the one shown in Fig. 1. In such images, the brighter particles carry larger forces [8, 16]. Furthermore, force chains tend to align preferentially along the principal stress axes [17]. It can be helpful to think of a force-chain network as the backbone of strong forces that span a system, providing support for both static [18] and dynamic [16] loading. However, weaker forces can also play a stabilizing role, much as guy-wires do on an aerial tower [19, 20].
It is also possible for sets of particles to cluster together into larger geographical domains, with potentially distinct properties, that can have weak structural boundaries between them [21]. At the largest scale, granular materials as a whole exhibit bulk properties, such as mechanical stability or instability in response to shear or compression [22]. All the aforementioned spatial scales are potentially relevant for understanding phenomena such as transmission of acoustic waves [23], thermal conductivity and heat transfer [24], electrical properties [25] and more. The time scales of interactions in granular materials are also important, and they can vary over many orders of magnitude. For example, in systems under compression, statistical fluctuations of grain displacements depend fundamentally on the length of the strain step (i.e., ‘increment’) over which one makes measurements, as fluctuations over short windows are consistent with anomalous diffusion and those over longer windows are consistent with Brownian behaviour [26].

The principled study of such diverse characteristics and organization in a single system can be very challenging, and the development of appropriate physical, mathematical, statistical and computational models is important to attain a mechanistic understanding of granular materials. Traditionally, it has been common to model granular materials using either particulate-based or continuum-based frameworks [20]. However, both of these approaches are often implicitly agnostic to intermediate-scale organization, which is important for understanding both static granular packings [27] as well as granular dynamics [28]. Recently, tools from network science [29, 30] and related mathematical subjects—which include approaches that can account explicitly for mesoscale structures [31–34]—have been used successfully to study properties of granular materials across multiple spatial and temporal scales. The most common representation of a network, an important idea for the study of complex systems of interacting entities [35], is as a graph [30]. A graph consists of a set of nodes (to represent the entities) and a set of edges, each of which represents an interaction between a pair of entities (or between an entity and itself). Increasingly, more complicated network representations (such as multilayer networks [36]) are
also employed. Moreover, there is also a growing recognition that it is important to consider the impact of other features, such as spatial embedding and other spatial effects [37], on network structure and dynamics, rather than taking an approach that promises that ‘one size fits all’. Network science offers methods for quantitatively probing and analysing large, interacting systems whose associated networks have heterogeneous patterns that defy explanations attained by considering exclusively all-to-all, regular, or lattice-like interactions [29].

There are several open problems in granular physics that may benefit from network-science approaches. In particular, because granular materials have multiple relevant length and time scales [16, 38–41], it can be challenging to model and quantify their structural organization, material properties and responses to external loads [42–46]. However, although complex, the pairwise inter-particle interactions that underlie and govern the structure and behaviour of granular systems (and other particulate matter) render them amenable to various network representations and network-based analyses. Smart and Ottino [47] were among the first to explicitly suggest and formalize the use of ideas from network science to begin to study some of the difficult questions in granular physics. In their article, they highlighted the ability of a network-based perspective to complement traditional methods for studying granular materials and to open new doors for the analysis of these complex systems. One place in which network analysis may be especially useful is in quantifying how local, pairwise interactions between particles in a granular packing yield organization on larger spatial scales (both mesoscale and system-level). For example, in sheared or compressed granular packings, such organization can manifest as force chains or other intermediate-sized sets of particles that together comprise a collective structure. Network science provides approaches to extract and quantitatively characterize heterogeneous architectures at microscale, mesoscale and macroscale sizes, and one can use these methods to understand important physical phenomena, including how such multiscale organization relates to bulk material properties or to spatial and temporal patterns of force transmission through a material.

Network-based approaches should also be able to provide new insights into the mechanisms that govern the dynamics of granular materials. For example, as we will discuss, network analysis can provide quantitative descriptions of how the structure of a dense granular material evolves as a system deforms under external loads (such as those induced by compression, shear, tapping or impact), and it can be helpful for describing certain aspects of complex dynamics (such as granular flows). Furthermore, it is important to consider the relative time scales of dynamics (e.g., sound propagation) on networks and the evolution of network structure. We also expect ideas from temporal networks [48] or adaptive networks [49] to be fruitful for studying faster structural dynamics, and investigation of granular dynamics in general should benefit from the development of both novel network representations and methods of network analysis that are designed specifically to understand temporally evolving systems. Another important problem in the study of granular materials is to predict when and where a granular system will fail. There has been some progress made in this area using network-based approaches, but it is important to continue to develop and apply tools from network analysis and related areas to gain a deeper understanding of which network features regulate or are most indicative of eventual failure. Another exciting direction for future work is to combine network-based approaches with questions about material design. In particular, can one use network-based approaches to help engineer granular systems—or other materials that are amenable to network representations—with desired and specialized properties?

It is also important to note that network-based representations and methods of analysis can provide insightful descriptions of granular materials with various additional complexities, such as systems that are composed of differently-shaped particles, three-dimensional (3D) materials and so on. This flexibility makes the application of tools from network science a powerful approach for studying the structural
properties and dynamics of granular systems. Such a framework also allows one to compare network architectures in diverse situations, such as between simulations and experiments, across systems that are composed of different types of particles or exposed to different loading conditions, and more. Exploiting these capabilities will yield improved understanding of which properties and behaviours of granular materials are general, versus which are specific to various details of a system. With the continued development of physically-informed network-analysis tools, network-based approaches show considerable promise for further development of both qualitative and quantitative descriptions of the organization and complex behaviour of granular materials.

The purpose of our article is to review the nascent application of network theory (and related topics) to the study of granular materials. We begin in Section 2.1 with a mathematical description of networks. In Section 2.2, we briefly review a set of measures that one can calculate on graphs and which have been useful in past investigations of granular materials. In Section 3, we review several ways in which granular materials have been represented as networks, and we discuss investigations of such networks to quantify heterogeneous, multiscale organization in granular materials and to understand how these systems evolve when exposed to external perturbations. We also point out insights into the underlying physics that have resulted from network-based investigations of granular matter. We close in Section 4 with some thoughts on the many remaining open questions, and we describe a few specific future directions that we feel are important to pursue. We hope that our review will be helpful for those interested in using tools from network science to better understand the physics of granular systems, and that it will spur interest in using these techniques to inform material design.

2. Network construction and characterization

2.1 What is a network?

It is often useful to model a complex system as a network, the simplest type of which is a graph [29, 50]. For most of our article, we will use the terms network and graph synonymously, but the former concept is more general than the latter. A graph \( G \) consists of nodes (i.e., vertices), where pairs of nodes are adjacent to each other via edges (i.e., links). We denote the set of nodes by \( V \) and the set of edges by \( E \).

A node can also be adjacent to itself via a self-edge (which is also sometimes called a self-loop), and a multi-edge describes the presence of two or more edges that are attached to (i.e., incident to) the same pair of nodes. (Unless we state otherwise, we henceforth assume that our networks have neither self-edges nor multi-edges.) The number of nodes in a graph is the size of the graph, and we also use the word ‘size’ in the same way for other sets of nodes. A subgraph of a graph \( G \) is a graph constructed using a subset of \( G \)’s nodes and edges.

An edge between two nodes represents some sort of relationship between them. For example, edges can represent information flow between different parts of the internet [53], friendship or other social interactions between people [54], trading between banks [55], anatomical or functional connections between large-scale brain regions [56, 57], physical connections between particles in contact [21, 58] and so on. Edges can be either unweighted or weighted, and they can be either undirected or directed [29]. In an unweighted (i.e., binary) network, an edge between two nodes is assigned a binary value (traditionally 0 or 1) to encode the absence or presence of a connection. In a weighted network, edges can take a variety

---

1 Indeed, it is increasingly important to examine network representations that are more complicated than graphs (see Sections 4.1 and 2.2.10)—such as multilayer networks [36], simplicial complexes [51] and others—and it is also essential to study dynamical processes on networks, rather than focusing exclusively on structural characteristics [52].
of different values to convey varying strengths of relationships between nodes. In an undirected network, all edges are bidirectional, so one assumes that all relationships are reciprocal. In a directed network, however, edges have a direction that encodes a connection from one node to another.

An adjacency matrix is a useful way to represent the information in a graph. For an unweighted and undirected graph, an element of the adjacency matrix $A$ of an $N$-node network is

$$A_{ij} = \begin{cases} 1, & \text{if there is an edge between nodes } i \text{ and } j, \\ 0, & \text{otherwise,} \end{cases}$$

(2.1)

where $i, j \in \{1, \ldots, N\}$. For a network in which nodes do not have labels, one can apply a permutation to $A$’s rows and columns—one uses the same permutation for each—to obtain another adjacency matrix that represents the same network. In this article, when we refer to ‘the’ adjacency matrix $A$ of a graph with unlabelled nodes, we mean any one of these matrices, which have the same properties (spectra, etc.).

For a weighted graph, if nodes $i$ and $j$ are adjacent via an edge, we denote the corresponding edge weight by $w_{ij}$ (which is usually given by a non-negative real number (e.g., the value of the normal or tangential component of the force between two contacting particles)). An element in the associated weighted adjacency matrix (which is sometimes called a weight matrix) $W$ is

$$W_{ij} = \begin{cases} w_{ij}, & \text{if there is an edge between nodes } i \text{ and } j, \\ 0, & \text{otherwise.} \end{cases}$$

(2.2)

For the more general case of a weighted, directed graph, if there is an edge from node $j$ to node $i$, then we let $w_{ij}$ represent the weight of that edge [29]. The associated weighted and directed adjacency matrix $W$ is

$$W_{ij} = \begin{cases} w_{ij}, & \text{if there is an edge from node } j \text{ to node } i, \\ 0, & \text{otherwise.} \end{cases}$$

(2.3)

An adjacency matrix associated with an undirected network is symmetric, but an adjacency matrix for a directed network need not be (and is symmetric if and only if all directed edges are reciprocated). In the present review, we primarily consider undirected networks, although we will occasionally make remarks about directed situations.

For weighted graphs, it is often also important to consider a binary adjacency matrix $A$ associated with a weight matrix $W$. Note that $A$ captures only the connectivity of nodes (i.e., their adjacencies), irrespective of how strongly they interact with each other. In terms of $W$, the corresponding binary network (which can be either directed or undirected) is

$$A_{ij} = \begin{cases} 1, & \text{if } W_{ij} \neq 0, \\ 0, & \text{otherwise.} \end{cases}$$

(2.4)

It is common to use terms like network topology when discussing structural properties of $A$, and sometimes one uses terms like network geometry when discussing properties that also depend on edge weights.

---

2 We do not consider edges with negative weights, although it may be interesting to do so in future work if there is an appropriate physical reason.
Fig. 2. From a packing to a network. (a) A sample packing of grains. (b) Representation of particles as network nodes. (c) Representation of contacts as network edges. (d) Graph representation of nodes and edges. The edges highlighted in green illustrate a cycle, which is a loop of physical contacts. The edges highlighted in peach illustrate a set of triangular loops, which are minimally-rigid structures in the context of structural rigidity. Edge weights often represent contact forces, which we illustrate here with different line widths. The degree $k$ of a node is equal to the number of edges attached (i.e., incident) to that node, and the strength (i.e., weighted degree) $s$ of a node is given by the sum of the weights of the edges attached to that node. One can (e) encode an unweighted (i.e., binary) graph as an unweighted adjacency matrix and (f) encode a weighted graph as a weighted adjacency matrix.

Because we will also employ ideas from subjects like algebraic topology (see Section 2.2.10), we will need to be very careful with such terminology.

The network representations that have been used to study granular matter (and other kinds of materials) employ diverse definitions of edges (both weighted and unweighted and both directed and undirected), and some generalizations of graphs have also been considered. See Fig. 2 for a schematic showing possible choices of nodes and edges for a network representation of a granular packing. A variety of tools and measures from network analysis have been used to study granular networks. We discuss some of these ideas in Section 2.2.

2.2 Some tools for characterizing granular networks

Network theory [29] provides myriad ways to characterize and quantify the topological and geometrical organization of complex networks. Thus, different network methods can reveal different important features of the underlying system, and these features, in turn, can help explain how a system behaves in certain situations. In the context of granular matter, for example, it is often desirable to understand the stability of a material, mechanical responses to external stresses, or wave propagation through a system. Recent investigations have demonstrated that network analysis can inform understanding of the
mechanisms that underlie these phenomena. In this section, we discuss several network concepts; and in Section 3, we describe how they have been used for the study of granular materials. We are, of course, not presenting an exhaustive list of methods from network science. See [29] and other books and reviews (and references therein) for discussions of other tools. For simplicity, we primarily give definitions for undirected networks, though many of the ideas that we present also have directed counterparts. We start with basic network diagnostics, and also discuss some more complicated methods.

2.2.1 Degree. One local property of a network is node degree. In an undirected network, a node’s degree is equal to the number of edges that are attached to it (see Fig. 2d). We denote the degree of node $i$ as $k_i$, and we recall that $N$ denotes the total number of nodes. For an unweighted graph with adjacency matrix $A$, one can calculate $k_i$ with the formula

$$k_i = \sum_{j=1}^{N} A_{ij}. \quad (2.5)$$

One can generalize the idea of degree to strength (i.e., weighted degree) using a weight matrix $W$ [59, 60]. The strength $s_i$ of node $i$ is equal to the sum of the weights of the edges that are attached to that node:

$$s_i = \sum_{j=1}^{N} W_{ij}. \quad (2.6)$$

Its associated degree $k_i$ is still given by Eq. (2.5).

A common network representation of granular materials is to treat particles as nodes and physical contacts between particles as either unweighted or weighted edges (see Fig. 2a–c). In this representation, node degree and node strength quantify information at the scale of single particles.

One can compute the mean degree (a global property) of a network by calculating

$$\langle k \rangle = \frac{1}{N} \sum_{i} k_i, \quad (2.7)$$

and one can similarly compute the mean strength with the formula

$$\langle s \rangle = \frac{1}{N} \sum_{i} s_i. \quad (2.8)$$

In an undirected network, mean degree is related to $N$ and the total number $m$ of edges through the relation $\langle k \rangle = \frac{2m}{N}$. It is sometimes useful to characterize a network using its degree distribution $P(k)$ [29], which gives the probability that a node has degree $k$.

When one represents a granular packing as a contact network (see Fig. 2 and Section 3.1), which is a binary network (i.e., unweighted network), the degree $k_i$ of node $i$ is known more commonly in the physics literature as its contact number or coordination number $Z_i$. If every node has the same degree (or even if most nodes have this degree), such as in a regular lattice, one often refers to the mean coordination number $Z$ of the lattice or packing. It is well-known that coordination number is related to the stability of granular systems [61] and plays a critical role in the jamming transition [62–64], a change of phase.
from an underconstrained state to a rigid state that is characterized by the onset of mechanical stability.\(^3\) We discuss these ideas further throughout the review.

2.2.2 Walks and paths. In a network, a \textit{walk} is an alternating sequence of nodes and edges that starts and ends at a node, such that consecutive edges are both incident to a common node. A walk thus describes a traversal from one node to another node (or to itself) along edges of a network. A \textit{path} is a walk that does not intersect itself or visit the same node (or the same edge) more than once, except for a \textit{closed path}, which starts and ends at the same node (see Section 2.2.3). One can compute the number of (unweighted) walks of a given length from a binary, undirected adjacency matrix \(A\) [29]. The length \(l\) of an unweighted walk is defined as the number of edges in the associated sequence (counting repeated edges the number of times that they appear). Letting \(\Xi^l_{ij}\) denote the number of walks of length \(l\) between nodes \(i\) and \(j\), one calculates

\[
\Xi^l_{ij} = [A^l]_{ij}. \tag{2.9}
\]

Various types of random walks yield short paths between nodes in a network, and such ideas (and their relation to topics such as spectral graph theory) are very insightful for studying networks [65].

In an undirected network, a path from node \(i\) to node \(j\) is necessarily also a path from node \(j\) to node \(i\). However, this is not typically true in directed networks, which have sometimes been utilized in studies of granular force chains (see e.g., [13]). Depending on a network’s structure, there may be several or no paths between a given pair of nodes. An undirected network is called \textit{connected} if there exists a path between each pair of nodes, and a directed network is called \textit{strongly connected} if there is a path between each pair of nodes [29]. (A directed network is called \textit{weakly connected} if its associated undirected network is connected, and a strongly connected network is necessarily also weakly connected.) In networks of granular packings, both the existence and lengths of paths can impact system behaviour. A connected network consists of a single \textit{component}. When a network has multiple components, it is common to study one component at a time (e.g., focusing on a \textit{largest connected component} (LCC), which is one that has the largest number of nodes).

The length of an unweighted path is the number of edges in the associated sequence, and it is sometimes also called the \textit{hop distance} (or occasionally, unfortunately, the \textit{topological distance}). Paths in a network can also be weighted by defining some (possibly abstract) notion of distance associated with the edges of the network. For example, in a spatially-embedded network [37], distance may refer to actual physical distance along an edge, in which case the length of a weighted path in the network is given by the sum of the physical distances along the sequence of edges in the path. However, one can also consider ‘distance’ more abstractly. For example, in a transportation or flow network, one can define a distance between two adjacent nodes to be some measure of resistance between those nodes, and then the length of a weighted path in such a network is given by the sum of the resistances along the sequence of edges in the path.\(^4\)

\(^3\) Unless we note otherwise, we use the phrase \textit{jamming} in the formal sense of the jamming transition as defined by [63, 64]. Packings of particles above the jamming point (a critical point related to the jamming transition) are rigid and overconstrained (i.e., ‘hyperstatic’), those at this point are marginally stable and exactly constrained (i.e., ‘isostatic’), and those below this point are underconstrained (i.e., ‘hypostatic’). Additionally, packings below the jamming point are sometimes called ‘un jammed’, and those above the jamming point are called ‘ jammed’.

\(^4\) One can also calculate distances between nodes if they occupy positions in a metric space (such as a latent one that determines the probability that a pair of nodes is adjacent to each other) [66], and the properties of that metric space can influence the distances (namely, the ones along a network) that concern us.
We use the term network distance to indicate a distance between two nodes (which can be either unweighted or weighted) that is computed by summing along edges in a path. A geodesic path—that is, a shortest path (which need not be unique)—between two nodes can be particularly relevant (though other short paths are often also important), and a breadth-first search (BFS) algorithm [67] is commonly employed to find geodesic paths in a network. The diameter of a graph is the maximum geodesic distance between any pair of nodes.

Denoting the shortest, unweighted network distance (i.e., shortest-path distance) between nodes \( i \) and \( j \) as \( d_{ij} \), the mean shortest-path distance \( L \) between pairs of nodes in a graph is [68]

\[
L = \frac{1}{N(N-1)} \sum_{i \neq j} d_{ij}.
\]

(2.10)

Note that one must be cautious when computing the mean shortest-path distance on disconnected networks (i.e., on networks that have more than one component), because the usual convention is to set the distance between two nodes in different components to be infinite [29]. Therefore, in a network with multiple components, the mean shortest-path distance \( L \) from Eq. (2.10) is infinite. One solution to this problem is to compute \( L \) for each component separately. Another network notion that relies on paths is network efficiency \([69, 70]\)

\[
E = \frac{1}{N(N-1)} \sum_{i \neq j} \frac{1}{d_{ij}}.
\]

(2.11)

One can generalize measures based on walks and paths to incorporate edge weights \([70, 71]\). Letting \( d_{ij}^w \) denote the shortest, weighted network distance between nodes \( i \) and \( j \), one can define a weighted mean shortest-path distance \( L^w \) and weighted efficiency \( E^w \) as in Eqs. (2.10) and (2.11), respectively, but now one uses \( d_{ij}^w \) instead of \( d_{ij} \). The network efficiency \( E \) is a normalized version of the Harary index of a graph [72]. Additionally, the convention \( d_{ij} = \infty \) (or \( d_{ij}^w = \infty \)) if there is no path from \( i \) to \( j \) allows one to use Eq. (2.11) (or its weighted counterpart \( E^w \)) on connected graphs or on graphs with more than one component. For both unweighted and weighted scenarios, large values of network efficiency tend to correspond to small values of mean shortest-path length, and vice versa. One can also readily generalize notions of paths, distances, and efficiency to directed networks \([69–71]\).

In later sections, we will describe the use of paths, walks and related ideas for investigating the structure of granular materials and their response to perturbations—including how these quantities change as a granular packing is compressed and goes through the jamming transition [73]—and we will also describe their use in specific applications, such as in understanding heat transfer through a granular material [24].

2.2.3 Cycles. A cycle (i.e., a closed walk) in a network is a walk that begins and ends at the same node [30]. As with other walks, one way to characterize a cycle is by calculating its length. An \( l \)-cycle is a cycle in which \( l \) edges are traversed (counting repeated edges the number of times that they appear in the cycle). A simple cycle is a cycle that does not include repeated nodes or edges, aside from one repetition of the origin node at the termination of a closed cycle. Thus, for example, a simple 3-cycle in an undirected network is a triangle. For the remainder of this review, we assume that cycles are simple cycles, unless we explicitly state otherwise. In the context of a granular packing, one can directly map particle contact loops—sets of physically-connected grains arranged in a circuit—to cycles in a corresponding graphical representation (see Fig. 2d). The length \( l \) is odd for an odd cycle and even for an even cycle.
We briefly note a few related concepts that are used to examine cycles in graphs because of their relevance to several network-based studies of granular materials. These are the notions of cycle space, cycle basis, and minimum cycle basis [30, 74]. The cycle space of an undirected graph is the set of all simple cycles in a graph along with all subgraphs that consist of unions of edge-disjoint simple cycles (i.e., they can share nodes but not edges) [75, 76]. A cycle basis is a minimal set of simple cycles such that any element of the cycle space can be written as a symmetric difference of cycles in the cycle basis [75]. Finally, for unweighted networks, a minimum cycle basis is a basis in which the total length of all cycles in the basis is minimal. For weighted networks, it is a basis in which the sum of the weights of all cycles in the basis is minimal.

Minimum cycle bases can provide useful information about the structure and organization of cycles in a network, so several algorithms have been developed to extract them (see, for example, [77, 78]). Once one has determined a minimum cycle basis, one can examine the distribution of cycle lengths or define measures to quantify the participation of different nodes in cycles of different lengths. For example, Walker et al. [79, 80] defined the concept of a cycle-participation vector $X^\text{cycle}_i = [x^0_i, x^3_i, \ldots, x^l_i]$ for each node $i$. The elements of this vector count the number of cycles of each length in which node $i$ participates. In this definition, $x^3_i$ is the number of 3-cycles in which node $i$ participates, $x^4_i$ is the number of 4-cycles in which node $i$ participates, and so on (up to cycles of length $l$). If node $i$ is not part of any cycle, then $x^0_i = 1$ and $x^j_i = 0$ for all $j \geq 3$; otherwise, $x^0_i = 0$.

One reason to examine cycles in granular networks [58, 73, 81–84] is that they can help characterize mesoscale structural features of a network. Cycles that are non-trivial involve more than a single node, but they do not typically embody global structures of a large network. This makes them appealing for studying network representations of granular materials, because mesoscale features seem to play a role in the behaviour of these systems [21]. Perhaps the most important motivation, however, is that cycles appear to be relevant for the stability and rigidity of a system. Specifically, in the context of structural rigidity theory, 3-cycles tend to be stabilizing structures that can maintain rigidity under applied forces [85], whereas 4-cycles can deform under applied forces (see Section 3.1.2). In Section 3, we discuss in more detail how cycles can help characterize granular systems.

### Cluster coefficients

Cluster coefficients are commonly-used diagnostics to measure the density of triangles either locally or globally in a network [29]. For an unweighted, undirected network, the local clustering coefficient $C_i$ is usually defined as the number of triangles involving node $i$ divided by the number of triples centred at node $i$ [68, 86]. A triple is a set of three nodes that can include either three edges (to form a 3-cycle) or just two of them. In terms of the adjacency matrix and node degree, the local clustering coefficient is

$$C_i = \frac{\sum_{hj} A_{hi}A_{hj}A_{ij}}{k_i(k_i - 1)}$$

for $k_i \geq 2$ (and $C_i = 0$ if $k_i \in \{0, 1\}$). One can then calculate a global clustering coefficient of a network as the mean of $C_i$ over all nodes:

$$C = \frac{1}{N} \sum_i^N C_i.$$
There is also another (and simpler) common way of defining a global clustering coefficient in a network that is particularly useful when trying to determine analytical approximations of expectations over ensembles of random graphs [29, 87].

The notion of a local clustering coefficient has also been extended to weighted networks in several ways [59, 88–90]. In one formulation [59], a local, weighted clustering coefficient $C_w^i$ is defined as

$$C_w^i = \frac{1}{s_i(k_i - 1)} \sum_{j,h} \frac{(W_{ij} + W_{ih})}{2A_{ij}A_{ih}A_{jh}}$$

(2.14)

for strength $s_i > 0$ and degree $k_i \geq 2$. The quantity $C_w^i = 0$ if either $s_i = 0$ (so that $k_i = 0$) or $k_i = 1$. Recall that $W$ and $A$ are, respectively, associated weighted and unweighted adjacency matrices. The mean of $C_w^i$ over all nodes gives a weighted clustering coefficient $C_w$ of a network. As we will discuss later (see Sections 3.1.3 and 3.2.4), clustering coefficients have been employed in several studies of granular materials. For example, they have been used to examine stability in granular networks [58, 73, 82–84]. See Fig. 3a for an example of the spatial distribution of a clustering coefficient in a granular packing.

2.2.5 Centrality measures. In network analysis, one often calculates centrality measures in efforts to quantify the importances of particular nodes, edges, or other structures in a network [29]. Different types of centralities characterize importance in different ways. The degree centrality (i.e., degree) of a node, for example, is simply the number of edges attached to it (see Section 2.2.1). A few other types of centrality that have been used to study granular materials are closeness centrality, node betweenness centrality, edge betweenness centrality, and subgraph centrality.

Notions of closeness centrality of a node measure how close that node is to other nodes in a network [91]. For a given node $i$, the most common notion of closeness is defined as the inverse of the sum
over the shortest-path lengths from node \( i \) to all other nodes \( j \) in a network. That is, node \( i \)'s closeness centrality is

\[
H_i = \frac{N - 1}{\sum_{j \neq i} d_{ij}}. \tag{2.15}
\]

Note that if we use the convention that the distance between two nodes in different components is infinite, then Eq. (2.15) only makes sense for connected networks. For any network with more than one component, Eq. (2.15) yields a closeness centrality of 0.

The geodesic node betweenness centrality of node \( i \) is the fraction of geodesic paths (either unweighted or weighted) between distinct nodes (not including \( i \)) that traverse node \( i \) [91]. Let \( \psi_{gh}(i) \) denote the number of geodesic paths from node \( g \) to node \( h \) that traverse node \( i \) (with \( i \not\in \{g, h\} \)), and let \( \psi_{gh} \) denote the total number of geodesic paths from node \( g \) to node \( h \). The geodesic node betweenness centrality of node \( i \) is then

\[
B_i = \sum_{g, h; g \neq h} \frac{\psi_{gh}(i)}{\psi_{gh}}, \quad i \not\in \{g, h\}. \tag{2.16}
\]

Geodesic node betweenness can probe the heterogeneity of force patterns in granular networks. See Fig. 3b for an example spatial distribution of a geodesic node betweenness centrality in an experimental granular packing. One can also compute a geodesic edge betweenness centrality by calculating the fraction of shortest paths (either unweighted or weighted) that traverse it [92]. Let \( \psi_{gh}(i, j) \) denote the number of geodesic paths from node \( g \) to node \( h \) that traverse the edge that is attached to node \( i \) and node \( j \), and let \( \psi_{gh} \) denote the total number of geodesic paths from node \( g \) to node \( h \). The geodesic edge betweenness centrality of this edge is then

\[
B_{ij} = \sum_{g, h; g \neq h} \frac{\psi_{gh}(i, j)}{\psi_{gh}}. \tag{2.17}
\]

Another measure of node importance is subgraph centrality \( Y \) [93, 94], which quantifies a node’s participation in closed walks of all lengths. Recall from Section 2.2.2 that one can write the number of length-\( l \) walks from node \( i \) to node \( j \) in terms of powers of the adjacency matrix \( A \). To calculate closed walks of length \( l \) that begin and end at node \( i \), we take \( i = j \) in Eq. (2.9). The subgraph centrality of node \( i \), with a specific choice for how much we downweight longer paths, is then given by

\[
Y_i = \sum_{l=0}^{\infty} \frac{[A^l]_{ii}}{l!}. \tag{2.18}
\]

Because shorter walks are weighted more strongly than longer walks in Eq. (2.18), they contribute more to the value of subgraph centrality (In other contexts, centrality measures based on walks have also been used to compare the spatial efficiencies of different networks, and such ideas are worth exploring in granular materials [72]). One can also express subgraph centrality in terms of the eigenvalues and eigenvectors of the adjacency matrix [93]. Let \( v^\alpha_i \) denote the \( i \)th component of the \( \alpha \)th eigenvector \( v^\alpha \) of
A, and let $\lambda_\alpha$ denote the corresponding $\alpha$th eigenvalue. One can then write

$$Y_i = \sum_{\alpha=1}^{N} (v_i^\alpha)^2 e^{i\lambda_\alpha}. \quad (2.19)$$

One can then calculate a mean subgraph centrality $Y$ by averaging $Y_i$ over the nodes in a network. In one study of granular materials [58], a subgraph centrality was examined for weighted networks by considering the eigenvalues and eigenvectors of the weight matrix $W$ in Eq. (2.19).

One can also compute network bipartivity $R$ [95] to quantify the contribution to mean subgraph centrality $Y$ from closed walks of even length. In particular, the network bipartivity $R_i$ of node $i$ is

$$R_i = \frac{Y_{\text{even}}}{Y_i}, \quad (2.20)$$

where $Y_{\text{even}}$ is the contribution to the sum in Eq. (2.18) from even values of $l$ (i.e., even-length closed walks). As with other node diagnostics, one can average bipartivity over all nodes in a network to obtain a global measure, which we denote by $R$.

In Section 3, we will discuss calculations of closeness, betweenness, and subgraph centralities in granular packings. Obviously, our discussion above does not give an exhaustive presentation of centrality measures, and other types of centralities have also been used in studies of granular materials (see for example, [21]).

### 2.2.6 Subgraphs, motifs and superfamilies

One can interpret the local clustering coefficient in Eq. (2.12) as a relationship between two small subgraphs: a triangle and a connected triple. Recall that a subgraph of a graph $G$ is a graph constructed using a subset of $G$’s nodes and edges. Conceptually, one can interpret small subgraphs as building blocks or subunits that together can be used to construct a network. For example, in a directed network, there exist three possible 2-node subgraphs (i.e., dyads): the dyad in which node $i$ is adjacent to node $j$ by a directed edge, the dyad in which node $j$ is adjacent to node $i$ by a directed edge, and the dyad in which both of these adjacencies exist. In a directed, unweighted graph, there are 13 different connected 3-node subgraphs [96] (see Fig. 4).
The term *motif* is sometimes used for a small subgraph that occurs often in a particular network or set of networks (typically relative to some null model, such as a randomly rewired network that preserves the original degree distribution) [96–99]. Borrowing terminology from genetics, these motifs appear to be *overexpressed* in a network (or set of networks). Unsurprisingly, the number of *n*-node subgraphs increases very steeply with *n*, so identifying subgraphs in large networks is computationally expensive, and many algorithms have been developed to estimate the number of subgraphs in an efficient (though approximate) way; see, for example, [96, 100–104]. In applying algorithms for motif counting to data, one seeks to identify subgraphs that are present more often than expected in some appropriate random-network null model.

The over-representation of a motif in a network is often interpreted as indicative of its playing a role in the function of that network (though one has to be cautious about drawing such conclusions). For example, 3-node motifs can form feedforward loops in which there are directed edges from node *i*₁ to node *i*₂, from node *i*₂ to node *i*₃ and from node *i*₃ to node *i*₁. The identification and characterization of motifs has yielded insights into the structure and function of a variety of systems, including food webs [105], gene-regulation networks of yeast [96], neuronal networks of the macaque monkey [106], and others. For different types of networks, one can also identify so-called *superfamilies*, which are sets of networks that have similar motif-frequency distributions [98]. There also exists a less-stringent definition of a superfamily; in this definition, one disregards whether a subgraph is a motif in the sense of it being more abundant than expected from some random-graph null model and instead considers a superfamily to be a set of networks that have the same rank-ordering of the number of *n*-node subgraphs for some fixed value of *n* [107]. In either case, one can examine different superfamilies to help understand the role that specific motifs (or subgraphs) or sets of motifs (or subgraphs) may have in potentially similar functions of networks in a given superfamily.

Subgraphs, motifs and superfamilies have been examined in several studies that applied network analysis to granular materials [80, 108–110]. They have revealed interesting insights into the deformation and reconfiguration that occurs in granular systems for different types of loading conditions and external perturbations. We discuss these ideas further in Sections 3.1.4 and 3.3.3.

2.2.7 Community structure. Many real-world networks have structure on intermediate scales (*mesoscales*) that can arise from particular organizations of nodes and edges [31–34, 111]. The most commonly-studied mesoscale network property is *community structure* [31, 32], which describes sets of nodes, called *communities*, that are densely (or strongly) interconnected to each other but only weakly connected to other dense sets of nodes. In other words, a community has many edges (or large total edge weight, in the case of weighted networks) between its own nodes, but the number and/or weight of edges between nodes in different communities is supposed to be small. Once one has detected communities in a network, one way to quantify their organization is to compute and/or average various network quantities over the nodes (or edges) within each community separately, rather than over the entire network. For example, one can compute the size (i.e., number of nodes) of a community, mean path lengths between nodes in a given community, or some other quantity to help characterize the architecture of different communities in a network. Studying community structure can reveal useful insights about granular systems, whose behaviour appears to be influenced by mesoscale network features [21, 27, 112–117].

Community structure and methods for detecting communities have been studied very extensively [31]. We will briefly discuss the method of *modularity maximization* [29, 118, 119], in which one optimizes an (occasionally infamous) objective function known as *modularity*, as this approach has been employed previously in several studies of granular materials (see, e.g., Section 3.2.2). However, myriad other
approaches exist for studying community structure in networks. These include stochastic block models (SBMs) and other methods for statistical inference (which are increasingly favored by many scholars) [111], approaches based on random walks (e.g., InfoMap [120]), various methods for detecting local community structure (see, e.g., [121, 122]), edge-based communities [123], and many others.

The goal of modularity maximization is to identify communities of nodes that are more densely (or more strongly) interconnected with other nodes in the same community than expected with respect to some null model. To do this, one maximizes a modularity objective function

\[ Q = \sum_{ij} [W_{ij} - \gamma P_{ij}] \delta(g_i, g_j), \]  

where \( g_i \) is the community assignment of node \( i \) and \( g_j \) is the community assignment of node \( j \), and where the Kronecker delta \( \delta(g_i, g_j) = 1 \) if \( g_i = g_j \) and \( \delta(g_i, g_j) = 0 \) otherwise. The quantity \( \gamma \) is a resolution parameter that adjusts the relative average sizes of communities [124, 125], where smaller values of \( \gamma \) favor larger communities and larger values of \( \gamma \) favor smaller communities [126]. The element \( P_{ij} \) is the expected weight of the edge between node \( i \) and node \( j \) under a specified null model. In many contexts, the most common choice is to determine the null-model matrix elements \( P_{ij} \) from the Newman–Girvan (NG) null model [118, 127, 128], for which

\[ P_{ij}^{NG} = \frac{s_i s_j}{2m}, \]  

where \( s_i = \sum_j W_{ij} \) is the strength (and, for unweighted networks, the degree \( k_i \)) of node \( i \) and \( m = \frac{1}{2} \sum_{ij} W_{ij} \) is the total edge weight (and, for unweighted networks, the total number of edges) in the network. There are several other null models, which are usually based on a random-graph model, and they can incorporate system features (such as spatial information) in various ways [129]. In the next part of this subsubsection, we discuss a physically-motivated null model that is particularly useful for studying granular force networks.

Maximizing \( Q \) is NP-hard [130], so it is necessary to use computational heuristics to identify near-optimal partitions of a network into communities of nodes [124]. Two well-known choices are the Louvain [131] and the Louvain-like [132] locally greedy algorithms, which begin by placing all nodes in their own community, and they then iteratively agglomerate nodes when the resulting partition increases modularity. Because of the extreme near degeneracy of the modularity landscape (a very large number of different partitions can have rather similar values of the scalar \( Q \)), it is often useful to run such an algorithm many times to construct an ensemble of partitions, over which one can average various properties to yield a ‘consensus’ description of community structure [126, 129, 133, 134].

**Physical considerations.** Community-detection tools, such as modularity maximization, have often been applied to social, biological, and other networks [31, 32]. In applying these techniques to granular materials, however, it is important to keep in mind that the organization of particulate systems (such as the arrangements of particles and forces in a material) is subject to significant spatial and physical constraints, which can severely impact the types of organization that can arise in a corresponding network representation of the material. When studying networks that are embedded in real space or constructed via some kind of physical relation between elements, it is often crucial to consider the spatial constraints—and, more generally, a system’s underlying physics—and their effects on network architecture [37]. Such considerations also impact how one should interpret network diagnostics such as path lengths and centrality measures, the null models that one uses in procedures such as modularity maximization, and so on.
NG null model was constructed to be appropriate for networks in which a connection between any pair of nodes is possible. Clearly, in granular materials—as in other spatially-embedded systems [37]—this assumption is unphysical and therefore problematic.

Bassett et al. [126] defined a null model that accounts explicitly for geographical (and hence spatial) constraints in granular materials, in which each particle can contact only its nearest neighbours [27]. In the context of granular networks with nodes representing particles and edges representing forces between those particles, the geographical null model $P$ in [126] has matrix elements

$$P_{ij} = \rho A_{ij},$$

where $\rho$ is the mean edge weight in the network and $A$ is the binary adjacency matrix of the network. In this particular application, $\rho = \bar{f} := \langle f_{ij} \rangle$ is the mean inter-particle force. As we illustrate in Fig. 5, modularity maximization with the geographical null model [Eq. (2.23)] produces different communities than modularity maximization with the NG null model [Eq. (2.22)] [27, 112, 113].

**Generalization of modularity maximization to multilayer networks.** Although studying community structure in a given granular packing can provide important insights, one is also typically interested in how such mesoscale structures reconfigure as a material experiences external perturbations, such as those from applied compression or shear. To examine these types of questions, one can optimize a multilayer generalization of modularity to study multilayer granular force networks in which each layer represents a network at a different step in the evolution of the system (for example, at different time steps or at different packing fractions) [113]. In Fig. 6, we show a schematic of a multilayer construction that has been employed in such investigations. See [36, 135] for reviews of multilayer networks (including generalizations of this construction).

One way to detect multilayer communities in a network is to use a generalization of modularity maximization [136], which was derived for multilayer networks with interlayer edges between nodes in different layers that represent the same entity. For simplicity, we suppose that all edges are bidirectional. One maximizes

$$Q_{\text{multi}} = \frac{1}{2\eta} \sum_{ijr} \left[ (W_{ijr} - \gamma_q \mathcal{P}_{ijr}) \delta_{qr} + \omega_{qpr} \delta_{ij} \right] \delta(g_{iq}, g_{jr}),$$

where $\rho$ is the mean edge weight in the network and $A$ is the binary adjacency matrix of the network. In this particular application, $\rho = \bar{f} := \langle f_{ij} \rangle$ is the mean inter-particle force. As we illustrate in Fig. 5, modularity maximization with the geographical null model [Eq. (2.23)] produces different communities than modularity maximization with the NG null model [Eq. (2.22)] [27, 112, 113].

**Fig. 5.** Modularity maximization with different null models can reveal distinct types of community structures in granular networks. In a network representation in which particles are represented by nodes and contact forces are represented by weighted edges, (a) using the NG null model helps uncover contiguous domains in a granular system, and (b) using a geographical null model on the same network helps detect chain-like structures that are reminiscent of force chains. In both panels, nodes (i.e., particles) of the same colour are assigned to the same community.
Fig. 6. A schematic of a multilayer network with layer-dependent community structure. In this example, each layer represents a static granular force network in which nodes (i.e., particles) are adjacent to one another via intralayer weighted edges (e.g., representing contact forces). Additionally, the same particle in consecutive layers is adjacent to itself via an interlayer edge of uniform weight $\omega$. For clarity, we only show two such couplings, but these interlayer edges exist for all particles (and between all consecutive layers). One can extract communities that change across layers—for example, if layers represent time, these are time-evolving communities—to study mesoscale organization in a granular system and to help understand how it reconfigures due to external loading (such as compression). In this schematic, we use different colours to label the particles that belong to different communities. Note that the same community can persist across several (or all) layers and reconfigure in terms of its constituent particles and the mean strength of its nodes. We reproduced this figure, with permission, from [113].

where $W_{ijq}$ is the $(i,j)$th component of the $q$th layer of the adjacency tensor $W$ [137] associated with the multilayer network, $\mathcal{P}_{ijq}$ is the $(i,j)$th component of the $q$th layer of the null-model tensor, $\gamma_q$ is a resolution parameter (sometimes called a structural resolution parameter) for layer $q$, and $\omega_{jqr}$ is the interlayer coupling between layers $q$ and $r$. (In the context of multilayer representations of temporal networks, if $\omega_{jqr} = \omega$ for all $j$, $q$ and $r$, one can interpret $\omega$ as a temporal resolution parameter.) More specifically, $\omega_{jqr}$ is the strength of the coupling that links node $j$ in layer $q$ to itself in layer $r$. (This type of interlayer edge, which occurs between nodes in different layers that represent the same entity, is called a diagonal edge [36].) The quantities $g_{iq}$ and $g_{jr}$, respectively, are the community assignments of node $i$ in layer $q$ and node $j$ in layer $r$. The intralayer strength of node $j$ in layer $q$ is $s_{jq} = \sum_i W_{ijq}$, and the interlayer strength of node $j$ in layer $q$ is $\zeta_{jq} = \sum_r \omega_{jqr}$, so the multilayer strength of node $j$ in layer $q$ is given by $\kappa_{jq} = s_{jq} + \zeta_{jq}$. Finally, the normalization factor $\eta = \frac{1}{2} \sum_{jq} \kappa_{jq}$ is the total strength of the adjacency tensor.$^5$

Maximizing multilayer modularity [Eq. (2.24)] allows one to examine phenomena such as evolving communities in time-dependent networks or communities that evolve with respect to some other parameter, and communities in networks with multiple types of edges. Capturing such behaviour has been useful in many applications, including financial markets [128], voting patterns [136], international relations [139], international migration [140], disease spreading [129], human brain dynamics [141–143] and more. In the context of granular matter, multilayer community detection allows one to examine changes in community structure of a force network, in which communities can either persist or reconfigure, with respect to both particle content and the mean strength of nodes inside a community, due to applied loads on a system.

---

$^5$ In the study of multilayer networks, it is common to use the term ‘tensor’ to refer to a multidimensional array [36] (as is common in some disciplines [138]), and proper tensorial structures have been explored briefly in adjacency tensors [137].
2.2.8 Flow networks. One can examine many natural and engineered systems—such as animal and plant vasculature, fungi, and urban transportation networks [144–150]—from the perspective of flow networks (which are often directed) that transport a load (of fluids, vehicles, and so on) along their edges. It is of considerable interest to examine how to optimize flow through a network [29, 151]. A well-known result from optimization theory is the maximum-flow–minimum-cut theorem [29, 151, 152]: for a suitable notion of flow and under suitable assumptions, the maximum flow that can pass from a source node to a sink node is given by the total weight of the edges in the minimum cut, which is the set of edges with smallest total weight that, when removed, disconnect the source and the sink. A related notion, which applies to networks in which there is some cost associated with transport along network edges, is that of maximum-flow–minimum-cost. In this context, one attempts to find a route through a network that maximizes flow transmission from source to sink, while minimizing the cost of flow along network edges [151, 152]. The maximum-flow–minimum-cut and maximum-flow–minimum-cost problems are usually examined under certain constraints, such as flow conservation at each node and an upper bound (e.g., limited by a capacitance) on flow through any edge. One can examine granular networks from such a perspective by considering a flow of force along a network formed by contacting grains. We discuss relevant studies in Section 3.3.1.

2.2.9 Connected components and percolation. Sometimes it is possible to break a network into connected subgraphs called components (which we introduced briefly in Section 2.2.2). A component, which is sometimes called a cluster, is a subgraph $G_C$ of a graph $G$ such that at least one path exists between each pair of nodes in $G_C$ [29]. Components are maximal subsets in the sense that the addition of another node of $G$ to a component destroys the property of connectedness. An undirected graph is connected when it consists of a single component. Networks with more than one component often have one component that has many more nodes than the others, so there can be one large component and many small components. One can find the components of a graph using a breadth-first search (BFS) algorithm [67], and one can determine the number of components by counting the number of 0 eigenvalues of a graph’s combinatorial Laplacian matrix [29]. To study graph components, one can also use methods from computational algebraic topology. Specifically, the zeroth Betti number $\beta_0$ indicates the number of connected components in a graph [153] (see Section 2.2.10).

Percolation theory [29, 154–156], which builds on ideas from subjects such as statistical physics and probability theory, is often used to understand the emergence and behaviour of connected components in a graph [52]. For example, in the traditional version of what is known as bond percolation (which is also traditionally studied on a lattice rather than on a more general network) [157], edges are occupied with probability $p$, and one examines quantities such as the size distributions of connected components as one varies the parameter $p$, called the bond occupation probability. It is especially interesting to determine a critical value $p_c$, called the percolation threshold, at which there is a phase transition: below $p_c$, there is no percolating component (or cluster), which spans the system and connects opposite sides; above $p_c$, there is such a cluster [157, 158]. In the latter scenario, it is common to say that there is a ‘percolating network’. In percolation on more general networks, one can study how the size of the largest component, as a fraction of the size of a network, changes with $p$. Related ideas also arise in the study of components in Erdős–Rényi random graphs $G(N, p)$, in which one considers an ensemble of $N$-node graphs and $p$ is the independent probability that an edge exists between any two nodes [29, 30, 159, 160]. In the limit $N \to \infty$, the size of the LCC undergoes a phase transition at a critical probability $p_c = 1/N$. When $p < p_c$, the ER graph in expectation does not have a giant connected component (GCC); at $p = p_c$, a GCC emerges whose size scales linearly with $N$ for $p > p_c$. Similarly, for bond percolation on networks, a transition occurs at a critical threshold $p_c$, such that for $p > p_c$, there is a GCC.
GRANULAR NETWORKS 505

(sometimes also called a ‘giant cluster’ or ‘percolating cluster’) whose size is a finite fraction of the total number \(N\) of nodes as \(N \to \infty\) [29, 52]. When studying percolation on networks, quantities of interest include the fraction of nodes in the LCC, the mean component size, the component-size distribution and critical exponents that govern how these quantities behave just above the percolation threshold [29, 158, 161].

We will see in Section 3 that it can be informative to use ideas from percolation theory to study the organization of granular networks. For example, it is particularly interesting to examine how quantities such as the number and size of connected components evolve with respect to packing density (or another experimental parameter). [28, 73, 162–169]. Some studies have considered connectivity percolation transitions, which are characterized by the appearance of a connected component that spans a system (i.e., a percolating cluster, as reflected by an associated GCC in the infinite-size limit of a network); or rigidity percolation transitions, which can be used to examine the transition to jamming [170–177].

Rigidity percolation is similar to ordinary bond percolation (which is sometimes used to study connectivity percolation), except that edges represent the presence of rigid bonds between network nodes [178, 179] and one examines the emergence of rigid clusters in the system with respect to the fraction of occupied bonds. One can also study percolation in force networks by investigating the formation of connected components and the emergence of a percolating cluster of contacts as one varies a force threshold, which is a threshold applied to a force-weighted adjacency matrix (representing contact forces between particles) to convert it to a binary adjacency matrix [73, 163–169, 176, 180–182]. (See Section 3.2.4 for additional discussion.) However, it is important to note that when studying networks of finite size, one needs to be careful with claims about GCCs and percolation phase transitions, which are well-defined mathematically only in the limit of infinite system size.

2.2.10 Methods from algebraic topology and computational topology. The tools that we have described thus far rely on the notion of a dyad (i.e., a 2-node subgraph) as the fundamental unit of interest (see Fig. 7a). However, recent work in algebraic topology and computational topology [153, 183–185] offers a complementary view, in which the fundamental building blocks that encode relationships between elements of a system are \(k\)-simplices (each composed of \(k + 1\) nodes), rather than simply nodes and dyadic relations between them (see Fig. 7b). These structures can encode ‘higher-order’ interactions and can be very useful for understanding the architecture and function of real-world networks (e.g., they yield a complementary way to examine mesoscale network features), and they have been insightful in the study of sensor networks [186], contagion spreading [187], protein interactions [188], neuronal networks [51, 189], and many other problems. See [190, 191] for further discussion and pointers to additional applications. The discussion in [192] is also useful.

A collection of simplices that are joined in a compatible way is called a simplicial complex, which is a generalization of a graph that can encode non-dyadic relations [185]. More precisely, and following [51], we define an (abstract) simplicial complex \(\mathcal{X}\) as a pair of sets: \(V_\mathcal{X}\), called the vertices (or nodes); and \(S_\mathcal{X}\), called the simplices, each of which is a finite subset of \(V_\mathcal{X}\), subject to the requirement that if \(\sigma \in S_\mathcal{X}\), then every subset \(\tau\) of \(\sigma\) is also an element of \(S_\mathcal{X}\). A simplex with \(k\) elements is called a \((k - 1)\)-simplex, and subsets \(\tau \subset \sigma\) are called faces of \(\sigma\). Using this notation, a 0-simplex is a node, a 1-simplex is an edge and its two incident nodes (i.e., a dyad), a 2-simplex is a filled triangle, and so on (see Fig. 7b). One type of simplicial complex that can be used to encode the information in a graph is a clique complex (sometimes also called a flag complex); we show an example in Fig. 8. To construct the clique complex of a graph \(G\), one associates every \(k\)-clique (a complete—i.e., fully connected—subgraph of \(k\) nodes) in \(G\) with a \((k - 1)\)-simplex. One can thus think of building the clique complex of a graph \(G\) as ‘filling in’ all of the \(k\)-cliques in \(G\) (see Fig. 7c). Note that we use the terms \(k\)-simplex and \(k\)-clique
Fig. 7. Algebraic topology and clique complexes. (a) An interaction in a graph is a dyad (i.e., a 2-node subgraph) or a node and a self-edge. (b) An alternative fundamental unit is a $k$-simplex. A 0-simplex is a node, a 1-simplex is an edge, a 2-simplex is a filled triangle, and so on. (c) A collection of $k$-simplices is called a simplicial complex, and one type of simplicial complex that can be used to encode the information in a graph is a clique complex (sometimes also called a flag complex). One constructs a clique complex by taking every $k$-clique (a complete subgraph of $k$ nodes) in a graph $G$ to be a simplex of the same number of nodes. (d) An interesting feature that can occur in a simplicial complex is a cycle, which is a closed arrangement of a collection of $k$-simplices. The purple edges in the upper object indicate a one-dimensional cycle that encloses a region filled in by simplices, whereas the purple edges in the lower object indicate a one-dimensional cycle that encloses a hole. (e) One can use a filtration to decompose a weighted graph into a sequence of binary graphs. For example, if one uses edge weight as a filtration parameter, one can represent a weighted graph as a sequence of unweighted graphs, which in turn yields a sequence of unweighted clique complexes. We adapted this figure, with permission, from [57].

because they are standard, but it is important not to confuse the use of $k$ in this context with the use of $k$ as the (also standard) notation for node degree.

One important feature of a simplicial complex is the potential presence of cycles. A cycle can consist of any number of nodes, and a $k$-dimensional cycle is defined as a closed arrangement of $k$-simplices, such that a cycle has an empty boundary. For example, Fig. 7d illustrates a closed arrangement of 1-simplices (i.e., edges) that forms a one-dimensional cycle. It important to distinguish between cycles that encircle a region that is filled by simplices with cycles that enclose a void (which is often called a ‘hole’ for the case of one-dimensional cycles). For example, the set of purple edges in the object in the upper portion of Fig. 7d constitutes a one-dimensional cycle that surrounds a region filled by 2-simplices (i.e., filled triangles), whereas the set of purple edges in the object in the lower portion of Fig. 7d constitutes a one-dimensional cycle that encloses a hole.

Characterizing the location and prevalence of void-enclosing cycles in the clique complex of a network representation of a granular packing can offer fascinating insights into the packing’s structure [163]. One way to do this is by computing topological invariants such as Betti numbers [153, 165, 183]. The $k$th

\[\text{filtration and persistent homology}\]

\[\text{parameter value}\]

---

6 Although we use the term cycle, which is standard in algebraic topology, note that this concept of a cycle is distinct from (though related to) the standard network-science use of the word ‘cycle’ (see Section 2.2.3). The latter is sometimes called a circuit, a term that we will use occasionally for clarity (especially given our focus on connected graphs).

7 The precise mathematical definition of a cycle requires a more detailed presentation than what we include in our present discussion. For more information and further details from a variety of perspectives, see [153, 183, 185, 188, 190, 192–194].
Betti number $\beta_k$ counts the number of inequivalent $k$-dimensional cycles that enclose a void, where two $k$-dimensional cycles are *equivalent* if they differ by a boundary of a collection of $(k+1)$-simplices. In other words, the $k$th Betti number $\beta_k$ counts the number of non-trivial *equivalence classes* of $k$-dimensional cycles and can thus also be interpreted as counting the number of voids (i.e., ‘holes’ of dimension $k$). The zeroth Betti number $\beta_0$ gives the number of connected components in a network, the first Betti number $\beta_1$ gives the number of inequivalent one-dimensional cycles that enclose a void (i.e., it indicates loops), the second Betti number $\beta_2$ gives the number of inequivalent 2D cycles that enclose a void (i.e., it indicates cavities), and so on.

Another useful way to examine the topological features that are determined by equivalence classes of $k$-dimensional cycles (i.e., components, loops, cavities, and so on) is to compute persistent homology (PH) of a network. For example, to compute PH for a weighted graph, one can first decompose it into a sequence of binary graphs. One way to do this is to begin with the empty graph and add one edge at a time in order of decreasing edge weights (see Fig. 7e). More formally and following [189], this process can translate information about edge weights into a sequence of binary graphs as an example of what is called a filtration [185, 190]. The sequence $G_0 \subset G_1 \subset \cdots \subset G_{|E|}$ of unweighted graphs begins with the empty graph $G_0$, and one adds a single edge at a time (or multiple edges, if some edges have the same weight) in order from largest edge weight to smallest edge weight. (One can also construct filtrations in other ways.) Constructing a sequence of unweighted graphs in turn yields a sequence of clique complexes [195], allowing one to examine equivalence classes of cycles as one varies the edge weight $\theta$ (or another filtration parameter). Important values of $\theta$ include the weight $\theta_{birth}$ associated with the first graph in which an equivalence class (i.e., a topological feature) occurs (i.e., its *birth* coordinate) and the edge weight $\theta_{death}$ associated with the first graph in which the feature disappears (i.e., its *death* coordinate), such as by being filled in with higher-dimensional simplices or by merging with an older feature. One potential marker of the relative importance of a particular feature (a component, a loop, and so on) in the clique complex is how long it persists, as quantified by its *lifetime* $\theta_{birth} - \theta_{death}$ (although short-lived features can also be meaningful [190, 192]). A large lifetime indicates robust features that persist over many values of a filtration parameter. Persistence diagrams (PDs) are one useful way to visualize the evolution of $k$-dimensional cycles with respect to a filtration parameter. PDs encode birth and death coordinates of features as a collection of persistence points $(\theta_{birth}, \theta_{death})$ in a planar region. One can construct a PD for each Betti number: a $\beta_0$ PD (denoted by PD$_0$) encodes the birth and death of components in a network, a $\beta_1$ PD (denoted by PD$_1$) encodes the birth and death of loops and so on.

To demonstrate some key aspects of a filtration, the birth and death of topological features, and PDs, we borrow and adapt an example from Kramár et al. [165]. Consider the small granular force network in Fig. 8a; the nodes represent particles in a 2D granular packing, and the coloured edges represent the magnitude of the inter-particle forces (of which there are four distinct values) between contacting particles. In a 2D system like this one, the only relevant Betti numbers are $\beta_0$ and $\beta_1$, as all others are 0. In Fig. 8b, we show the *flag complex* (which is essentially the same as a *clique complex* [190]) of the granular network, where the colour of a triangle indicates the value corresponding to the minimum force along any of its edges. Computing PH on a flag complex (which has been done in several studies of PH in granular force networks [164–169]) only counts loops that include four or more particles. That is, it does not count 3-particle loops (which are sometimes called ‘triangular loops’). Loops with four or more particles are associated with *defects*, as they would not exist in a collection of monosized disks that
Fig. 8. An example force network, an associated filtration over the flag complex (i.e., clique complex), and PDs. (a) In the force network, colored edges represent the magnitude of the force between contacting particles, which are represented as nodes in the network. In order from smallest to largest, the four values of the force are $\theta_1$ (dark blue), $\theta_2$ (cyan), $\theta_3$ (green), and $\theta_4$ (red). (b) The flag complex is formed by filling in all 3-particle loops (i.e., triangular loops) with the smallest value of the force along any of its edges. Defining a filtration over the flag complex avoids counting these 3-particle loops. (c–f) The sequence of complexes corresponding to the filtration over the flag complex; one obtains the sequence by descending the four levels of the force threshold $\theta$. (g) The $\beta_0$ persistence diagram PD$\beta_0$. (h) The $\beta_1$ persistence diagram PD$\beta_1$. We adapted this figure, with permission, from [165].
are packed perfectly and as densely as possible into a ‘crystalline’ structure (which has only triangular loops) [165].

In Fig. 8c–f, we show the sequence of complexes that correspond to the filtration over the flag complex. One descends the four threshold levels (i.e., edge weights), beginning with the largest ($\theta_4$) and ending with the smallest ($\theta_1$). In Fig. 8g and h, we show the corresponding PDs for $\beta_0$ and $\beta_1$. It is helpful to discuss a few features of these diagrams. In PD$_0$, we observe four points that are born at $\theta_4$; these points correspond to the four connected components that emerge at the first level of the filtration in Fig. 8c. Two of the components merge into one component at $\theta_3$ (see Fig. 8d); this corresponds to the point at $(\theta_4, \theta_3)$. A new component forms at $\theta_3$ and dies at $\theta_2$; this is represented by the point at $(\theta_3, \theta_2)$ (see Fig. 8d). Additionally, two components born at $\theta_4$ die at $\theta_2$, corresponding to the two points at $(\theta_4, \theta_2)$. One can continue this process until the end of the filtration, where there is just a single connected component (see Fig. 8f). This component is born at $\theta_4$; it persists for all thresholds, and we use Kramár et al. [165]’s convention to give it a death coordinate of $-1$; this yields the persistence point at $(\theta_4, -1)$. In PD$_1$, we observe that a loop emerges at $\theta_3$ (see Fig. 8d), and it is then filled by triangles at $\theta_2$ (see Fig. 8e), leading to the point at $(\theta_3, \theta_2)$. Three more loops are born at $\theta_2$ and never die (see Fig. 8e); again using the convention in [165], we assign these features a death coordinate of 0, so there are three persistence points at $(\theta_2, 0)$. Finally, one more loop appears at $\theta_1$ and does not die (see Fig. 8e); this is represented by a point at $(\theta_1, 0)$.

Kramár et al. [165] gave an in-depth exposition of how to apply PH to granular networks, and we refer interested readers to this article for more information. Because studying PH is a general mathematical approach, it can be applied to different variations of force networks and can also be used on networks constructed from different types of experimental data (e.g., digital image data, particle-position data or particle-interaction data). Kramár et al. [165] also discussed a set of measures that can be used to compare and contrast the homology of force networks both within a single system (i.e., at two different packing fractions) and across different systems (e.g., if one uses particles of different sizes or shapes), and they explored the robustness of PH computations to noise and numerical errors. In Section 3.2.5, we further discuss applications of methods from algebraic and computational topology to granular materials.

2.3 Some considerations when using network-based methods

Because there are many methods that one can use to analyse granular networks and many quantities that one can compute to measure properties of these networks, it is useful to discuss some relationships, similarities and distinctions between them. Naturally, the meaning of any given network feature depends on how the network itself is defined, so we focus the present discussion on the most common representation of a granular system as a network. (See Section 3.3 for discussions of other representations.) In this representation (see Fig. 2), nodes correspond to particles and edges correspond to contacts between particles. Additionally, edge weights can represent quantities such as normal or tangential forces between particles. In this type of granular network, it is important to be aware of which network quantities explicitly take into account spatial information or physical constraints in a system, which consider only network topology, and which consider only network geometry (i.e., both topology and edge weights, but not other information). Granular materials have a physical nature and are embedded in real space, so such considerations are extremely important. For discussions of how these issues manifest in spatial networks more generally, see [37].

One way to explicitly include spatial or physical information into network analysis is to calculate quantities that are defined from some kind of distance (e.g., a Euclidean distance between nodes), whether directly or through a latent metric space, rather than a hop distance. For example, as discussed in Section 2.2.2, one can define the edge length between two adjacent nodes from the physical distance
between them, which allows quantities such as mean shortest path length, efficiency and some central-
ity measures to directly incorporate spatial information. However, traditional network features such as
degree and clustering coefficient depend only on network connectivity, although their values are influ-
enced by spatial effects. In Section 2.2.7, we also saw that one can incorporate physical constraints from
granular networks into community-detection methods by using a geographical null model, rather than
the traditional NG null model, in modularity maximization.

Different computations in network analysis can also probe different spatial, topological or geometri-
cal scales. For example, measures such as degree, strength and clustering coefficients are local measures
that quantify information about the immediate neighbourhood of a node. However, measures such as the
mean shortest path length and global efficiency are global in nature, as they probe large-scale network
organization. In between these extremes are mesoscale structures. A network-based framework can be
very helpful for probing various types of intermediate-scale structures, ranging from very small ones (e.g.,
motifs, such as small cycles) to larger ones (e.g., communities) and tools such as PH were designed to
reveal robust structural features across multiple scales. Crucially, although there are some clear qualitative
similarities and differences between various network-analysis tools (and there are some known quantita-
tive relationships between some of them [29]), it is a major open issue to achieve a precise understanding
of the relationships between different network computations. Moreover, in spatially-embedded systems
(as in any situation where there are additional constraints) one can also expect some ordinarily distinct
quantities to become more closely related to each other [37]. Furthermore, the fact that a granular particle
occupies a volume in space (volume exclusion) gives constraints beyond what arises from embeddedness
in a low-dimensional space.

3. Granular materials as networks

We now review network-based models and approaches for studying granular materials. Over the past
decade, network analysis has provided a novel view of the structure and dynamics of granular systems,
insightfully complementing and extending traditional perspectives. See [1–5] for reviews of non-network
approaches.

Perhaps the greatest advantages of using network representations and associated tools are their natural
ability to (1) capture and quantify the complex and intrinsic heterogeneity that manifests in granular
materials (e.g., in the form of force chains) and to (2) systematically and quantitatively investigate how the
structure and organization of a granular system changes when subjected to external loads or perturbations
(such as compression, shear, or tapping). In particular, network science and related subjects provide a set of
tools that help quantify structure (and changes in structure) over a range of scales—including local, direct
interactions between neighbouring particles; larger, mesoscale collections of particles that can interact and
reconfigure via more complicated patterns; and system-wide measurements of material (re)organization. It
is thought that local, intermediate, and system-wide scales are all important for regulating emergent, bulk
properties of granular systems, but it can be difficult to obtain a holistic, multiscale understanding of these
materials. For example, particle-level approaches may not take into account collective organization that
occurs on slightly larger scales, and continuum models and approaches that rely on averaging techniques
may be insensitive to interesting and important material inhomogeneities [42–44, 46].

Network representations also provide a flexible medium for modeling different types of granular
materials (and other particulate matter). For example, network analysis is useful for both simulation
and experimental data of granular materials, and methods from complex systems and network science
can help improve understanding of both dense, quasistatically-deforming materials as well as granular
flows. In any of these cases, one often seeks to understand how a system evolves over the course of
an experiment or simulation. To study such dynamics, one can examine a network representation of a system as one varies a relevant physical quantity that parameterizes the system evolution. For example, for a granular system in which the packing fraction increases in small steps as the material is compressed quasistatically, one can extract a network representation of the system at each packing fraction during the compression process and then study how various features of that network representation change as the packing fraction increases. Even a particular type of granular system is amenable to multiple types of network representations, which can probe different aspects of the material and how it evolves under externally applied loads. For instance, one can build networks based only on knowledge of the locations of particles (which, in some cases, may be the only information available) or by considering the presence or absence of physical contacts between particles.

If one knows additional information about the elements in a system or about their interactions, one can construct more complicated network representations of it. For example, it has long been known that granular materials exhibit highly heterogeneous patterns of force transmission, with a small subset of the particles carrying a majority of the load along force chains [196, 197]. Recall from Section 1 that, broadly speaking, a force chain (which is also sometimes called a force network) is a set of contacts that carry a load that is larger than the mean load [8, 16], and the mean orientation of a force chain often encodes the direction of the applied stress [17]. We illustrated an example of force-chain structure in Fig. 1, and we further discuss force-chain organization in Section 3.2. Because of the nature of the distribution of force values and the interesting way in which forces are spatially distributed in a material, it is often very useful to consider network representations of granular materials that take into account information about inter-particle forces (see Section 3.2) and to use network-based methods that allow one to quantitatively investigate how the structure of a force network changes when one includes only contacts that carry at least some threshold force (see Sections 3.2.4 and 3.2.5).

In our ensuing discussion, we describe several network constructions that have been used to study granular materials, indicate how they have been investigated using many of the concepts and diagnostics introduced in Section 2.2, and review how these studies have improved scientific understanding of the underlying, complex physics of granular systems.

### 3.1 Contact networks

A contact network is perhaps the simplest way to represent a granular system. Such networks (as well as the term ‘contact network’) were used to describe granular packings long before explicitly network science-based approaches were employed to study granular materials; see, for example, [198, 199]. The structure of a contact network encodes important information about a material’s mechanical properties. As its name suggests, a contact network embodies the physical connectivity and contact structure of the particles in a packing (see Fig. 2). In graph-theoretic terms, each particle in the packing is represented as a node, and an edge exists between any two particles that are in physical contact with one another. Note that it may not always be possible to experimentally determine which particles are in physical contact, and one may need to approximate contacts between particles using information about particle positions, radii, and inter-particle distances. (See Section 3.5 for details.) By definition (and however it is constructed), a contact network is unweighted and undirected, and it can thus be described with an unweighted and undirected adjacency matrix (see Section 2.1):

\[
A_{ij} = \begin{cases} 
1, & \text{if particles } i \text{ and } j \text{ are in contact}, \\
0, & \text{otherwise}. 
\end{cases} \quad (3.1)
\]
Fig. 9. An example of a contact network from a DEM simulation of a densely-packed 2D system of polydisperse, disk-shaped particles. In this case, the granular material was subjected to biaxial compression and constrained to move along a plane. This snapshot corresponds to a network at an axial strain before full shear-band formation. We adapted this figure, with permission, from [58].

Because the organization of a contact network depends on and is constrained by the radii of the particles and their locations in Euclidean space, a contact network is a \textit{spatially-embedded} graph [37]. In Section 3.2.2, we will see that this embedding in physical space has important consequences for the extraction of force-chain structures via community-detection techniques (see Section 2.2.7). In Fig. 9, we show an example of a contact network generated from a discrete element method (DEM) simulation (see Section 3.5) of biaxial compression [200]. The granular system in this figure is \textit{polydisperse}, as it has more than two types of particles. (In this case, the particles have different sizes.) If all particles are identical in a granular system, it is called \textit{monodisperse}; if there are two types of particles in a system, it is called \textit{bidisperse}. In practice, although the presence or absence of a contact is definitive only in computer simulations, one can set reasonable thresholds and perform similar measurements in experiments [201] (see Section 3.5). It is also important to note that packing geometry and the resulting contact network do not completely define a granular system on their own. In particular, one can associate a given geometrical arrangement of particles with several configurations of inter-particle forces that satisfy force and torque balance constraints and the boundary conditions of a system [202–205]. This is a crucial concept to keep in mind when conducting investigations based only on contact networks, and it also motivates the inclusion of contact forces to construct more complete network representations of granular systems (see Section 3.2).

In the remainder of this subsection, we review some of the network-based approaches for characterizing contact networks of granular materials and how these approaches have been used to help understand the physical behaviour of granular matter. We primarily label the following subsubsections according to the type of employed methodology. However, we also include some subsubsections about specific applications to certain systems.
3.1.1 Coordination number and node degree. One can study a contact network in several ways to investigate different features of a granular system. We begin our discussion by associating the mean node degree of a contact network with the familiar and well-studied coordination number (i.e., contact number) \( Z \). Although early investigations of granular materials did not consciously make this connection, the mean degree and coordination number are synonymous quantities. The contact degree \( k_i \) of particle \( i \) is the number of particles with which \( i \) is directly in contact, and one can calculate it easily from an adjacency matrix [see Eq. (2.5)]. A contact network is undirected, so its adjacency matrix \( A \) is symmetric, and its row sum and column sum each yield a vector of node degrees. The mean degree \( \langle k \rangle \) of a contact network [see Eq. (2.7)] is then the mean coordination number (i.e., contact number) \( Z \), and it gives the mean number of contacts per particle. As we noted previously (see Section 2.2.1), \( Z \) is an important quantity in granular systems because of its connection with mechanical stability and rigidity—which, loosely speaking, is the ability of a system to withstand deformations—and its characterization of the jamming transition \[63, 206\] and other mechanical properties. In particular, the condition for mechanical stability—that is, the condition to have all translational and rotational degrees of freedom constrained so that there is force and torque balance—in a packing of frictionless spheres in \( d \) dimensions \[61, 63, 64, 206, 207\] is

\[
Z \geq 2d \equiv Z_{iso}. \tag{3.2}
\]

The isostatic number \( Z_{iso} \) indicates the condition for isostaticity, which is defined as the minimum contact number that is needed for mechanical stability. One can use the coordination number (which is often tuned by changing the packing fraction \( \phi \)) as an order parameter to describe the jamming transition for frictionless spheres in 2D and 3D \[63, 64, 207\]. Specifically, there is a critical packing fraction \( \phi_c \) such that below \( \phi_c \), the contact number for these systems is \( Z = 0 \) (i.e. there are no load-bearing contacts), and at the critical packing fraction \( \phi_c \), the contact number jumps to the critical value \( Z_c = Z_{iso} = 2d \).

One can also generalize the use of the coordination number in order to examine mechanical stability and jamming in granular systems of frictional spheres. In these systems, the condition for stability is

\[
Z \geq Z_{iso}^m, \quad Z_{iso}^m \equiv (d + 1) + \frac{2N_m}{d}, \tag{3.3}
\]

where \( N_m \) is the mean number of contacts per particle that have tangential forces \( f_t \) equal to the so-called Coulomb threshold—that is, \( N_m \) is the mean number of contacts per particle with \( f_t = \mu f_n \), where \( \mu \) is the coefficient of friction and \( f_n \) is the normal force \[64, 207, 208\]—and \( Z_{iso}^m \) again designates the condition for isostaticity. Results from experimental systems have demonstrated that contact number also characterizes the jamming transition in frictional, photoelastic disks \[201\].

The coordination number has been studied for several years in the context of granular materials and jamming, and it is fruitful to connect it directly with ideas from network science. Several recent studies have formalized the notion of a contact network, and they deliberately modeled granular systems as such networks to take advantage of tools like those described in Section 2.2. Such investigations of contact networks allow one to go beyond the coordination number and further probe the rich behaviour and properties of granular materials—including stability and the jamming transition \[63\], force chains \[8, 9, 16\], and acoustic propagation \[21, 23, 42, 45\].

Perhaps the simplest expansion of investigations into the role of the coordination number is the study of the degree distribution \( P(k) \) of the contact network of a packing. Calculating degree distributions can provide potential insights into possible generative mechanisms of a graph \[29, 158\], although one has
to be very careful to avoid over-interpreting the results of such calculations [209]. In granular physics, it has been observed that the degree distribution of a contact network can track changes in network topology past the jamming transition in isotropically compressed simulations of a 2D granular system [73]. Specifically, the peak of $P(k)$ shifts from a lower value of $k$ to a higher value of $k$ near the transition. Moreover, changes in the mean degree $\langle k \rangle$ and its standard deviation can anticipate the onset of different stages of deformation in DEM simulations (i.e., molecular-dynamics simulations) of granular systems under various biaxial compression tests [58, 210].

3.1.2 Investigating rigidity of a granular system using a contact network. An important area of research in granular materials revolves around attempts to (1) understand how different types of systems respond when perturbed, and (2) determine what features of a system improve its integrity under such perturbations. As we noted in Section 3.1.1, it is well-known that coordination number (and hence node degree) is a key quantity for determining mechanical stability and understanding jamming in granular materials. However, contact networks obviously have many other structural features, and examining them can be very helpful for providing a more complete picture of these systems.

To the best of our knowledge, the stability of granular materials was first studied from a graph-theoretic standpoint in the context of structural rigidity theory [172, 179, 211], which has also been applied to amorphous solids more generally [62]. In structural rigidity theory, thought to have been studied originally by Maxwell [212], rods of fixed length are connected to one another by hinges, and one considers the conditions under which the associated structural graphs are able to resist deformations and support applied loads (see Fig. 10). A network is said to be minimally rigid (or isostatic) when it has exactly the number of bars needed for rigidity. This occurs when the number of constraints is equal to the number of degrees of freedom in the system (i.e., when Laman’s minimal-rigidity criterion is satisfied). The network is flexible if there are too few rods, and it is overconstrained (i.e., self-stressed) if there are more rods than needed for minimal rigidity. Triangles are the smallest isostatic structures in two dimensions [213–215]; there are no allowed motions of these structures that preserve the lengths and connectivity of the bars, so triangles (i.e., 3-cycles) do not continuously deform due to an applied force. In comparison, 4-cycles are structurally flexible and can continuously deform from one configuration to another while preserving the lengths and connectivity of the rods (see Fig. 10).

Extending a traditional network of rods and hinges, concepts from structural rigidity theory yield interesting insights into contact networks of particulate matter. See [216] for a discussion of some of
the earliest applications of such ideas to disordered systems and granular materials. Moukarzel [217] used structural rigidity theory to derive conditions for the isostaticity of a granular packing, and he tied the fact that random packings of particles tend to be isostatic to the origin of instabilities in granular piles. Later, similar concepts were used to show that in granular networks, cycles with an even number of edges allow contacting grains to roll without slipping when subject to shear; however, these relative rotations are ‘frustrated’ in cycles with an odd number of edges, so such cycles can act as stabilizing structures in a network [85]. Several later studies (such as [58, 73, 79–81, 83, 84, 210, 218–220]) have confirmed that contact loops are often stabilizing mesoscale features in a contact network of a granular material. We specifically consider the role of cycles in granular contact networks in Section 3.1.3.

Another type of network approach for understanding rigidity in granular systems is rigidity percolation [178, 179, 211] (see Section 2.2.9). Feng [170] conducted an early investigation of an idealized version of bond percolation in a granular context. It is now known that hallmarks of this bond-percolation transition occur below isostaticity: Shen et al. [175] identified that a percolating (i.e., system-spanning) cluster of non-load-bearing contacts forms at a packing density below the jamming point. One can also use a rigidity-percolation approach to examine if a network is both percolating and rigid (see Section 2.2.9). Note that a rigid granular network is also percolating, but a percolating network need not be rigid. Rigidity percolation relies on tabulating local constraints via a pebble game [172], which reveals connected, rigid regions (sometimes called ‘rigid clusters’) in a network. In a series of articles [221–224] on simulated packings, Schwarz and co-workers went beyond Laman’s minimal-rigidity criterion to investigate local versus global rigidity in a network, the size distribution of rigid clusters, the important role of spatial correlations, and the necessity of force balance. Building on the above work, [177] recently utilized a rigidity-percolation approach to identify floppy versus rigid regions in slowly-sheared granular materials and to characterize the nature of the phase transition from an underconstrained system to a rigid network. See also the recent article [225].

3.1.3 Exploring the role of cycles. We now consider the role of circuits (i.e., the conventional network notion of cycles, which we discussed in Section 2.2.3) in granular contact networks. Cycles in a contact network can play crucial stabilizing roles in several situations. Specifically, as we will discuss in detail in this section, simulations (and some experiments) suggest that (1) odd cycles (especially 3-cycles) can provide stability to granular materials by frustrating rotation among grains and by providing lateral support to surrounding particles, and that (2) a contact network loses these stabilizing structures as the corresponding granular system approaches failure.

Noting that 3-cycles are the smallest arrangement of particles that can support (via force balance) a variety of 2D perturbations to a compressive load without deforming the contact structure, Smart and Ottino [81] studied the effects of friction and tilting on the evolution of contact-loop organization in a granular bed. In their simulations, they implemented tilting by incrementally increasing the angle of a gravity vector with respect to the vertical direction, while preserving the orientation of the granular bed and maintaining quasistatic conditions. In untilted granular packings, they observed that lowering inter-particle friction yields networks with a higher density of 3-cycles and 4-cycles, where they defined the ‘density’ of an $l$-cycle to be the number of $l$-cycles divided by the total number of particles. By examining the contact network as they varied tilting angle, Smart and Ottino [81] also observed that the density of 4-cycles increases prior to failure—likely due to the fracture of stabilizing 3-cycles—and that this trend was distinguishable from changes in the coordination number alone.

Cycles have also been studied in the context of DEM simulations of dense, 2D granular assemblies subject to quasistatic, biaxial compression [58, 84, 210, 218]. In many of these studies, the setup consists
of a collection of disks in 2D that are compressed slowly at a constant strain rate in the vertical direction, while allowed to expand under constant confining pressure in the horizontal direction [58, 84, 218]. In another variation of boundary-driven biaxial compression, a sample can be compressed with a constant volume but a varying confining pressure [210]. Before describing specifics of the network analysis for these systems, it is important to note that for the previously described conditions, the axial strain increases in small increments (i.e., ‘steps’) as compression proceeds, and one can extract the inter-particle contacts and forces at each strain value during loading to examine the evolution of a system with respect to strain. Additionally, these systems undergo a change in behaviour from a solid-like state to a liquid-like state, and they are characterized by different regimes of deformation as one increases axial strain [200]. In particular, the granular material first undergoes a period of strain hardening, followed by strain softening, after which it enters a critical state. In the strain-hardening regime, the system is stable and the shear stress increases monotonically with axial strain up to a peak value. After the peak shear stress, strain softening sets in; this state is marked by a series of steep drops in the shear stress that indicate reduced load-carrying capacity. Finally, in the critical state, a persistent shear band has fully formed, and the shear stress fluctuates around a steady-state value. The shear band is a region of localized deformation and provides one signature of material failure [226]. Inside the shear band, force chains can both form and buckle [227]. One can also associate increases in the energy dissipation rate of the system with particle rearrangements (such as those that occur during force-chain buckling) and loss of stability in the material.

Examining the temporal evolution of cycles in such an evolving granular contact network can reveal important information about changes that occur in a material during deformation. Using DEM simulations (see the previous paragraph), Tordesillas et al. [84] computed the total number of cycles of different lengths in a minimal cycle basis (see Section 2.2.3) of a contact network at each strain state during loading, and they observed that there are many more 3-cycles and 4-cycles than longer cycles in the initial, solid-like state of the system. However, as axial strain increases and one approaches the maximum shear stress, the total number of 3-cycles falls off steeply. (The same is true for 4-cycles, though the effect is less dramatic.) Additionally, during axial-strain steps (i.e., axial-strain ‘intervals’) corresponding to drops in shear stress, Tordesillas et al. [84] observed large increases in the number of 3-cycles and 4-cycles that open up to become longer cycles. In Fig. 11a, we show an example of the evolution of cycle organization with increasing axial strain for a subset of particles from a DEM simulation of a granular material under biaxial compression carried out by Walker and Tordesillas [58]. The authors observed that in this system, both the global clustering coefficient $C$ [see Eq. (2.13)] and the mean subgraph centrality $Y$ [see Eq. (2.19)] decrease with increasing axial strain, drop sharply at peak shear stress, and then level out (see Fig. 11b and c). Recalling that $C$ is a measure of triangle density in a graph and that subgraph centrality measures participation of nodes in closed walks (with more weight given to shorter paths), these results also imply that the loss of small cycles co-occurs with the deformation and failure of a system due to increasing load. Walker and Tordesillas [58] also computed the network bipartivity $R$ [95] of the contact network to quantify the contribution to mean subgraph centrality $Y$ from closed walks of even length [see Eq. 2.20]. They observed that $R$ increases with increasing axial strain, revealing that closed walks of even length become more prevalent during loading (see Fig. 11c). The authors suggested that this trend may be due to a decrease in the prevalence of 3-cycles (which are stabilizing, as discussed in Section 2.2.3 and elsewhere). Tordesillas et al. [218] also examined the stability of cycles of various lengths in both DEM simulations and experimental data, and they observed that, during loading, 3-cycles tend to be more stable (as quantified by a measure of stability based on a structural-mechanics framework [228]) than cycles of other lengths in a minimal cycle basis of the network.

Minimal cycle bases and the easier-to-compute subgraph centrality have also been used to examine fluctuations in kinetic energy in simulations of deforming sand. Walker et al. [79] computed a minimal
Fig. 11. Evolution of cycles in a deforming granular material. (a) One can track a subset of particles and their corresponding contact network from a DEM simulation for increasing axial strain values $|\varepsilon_2|$. The system consists of 5098 spherical, polydisperse particles that were subjected to a quasistatic, biaxial compression test. At the smallest displayed axial strain, the set of particles in this figure yields a network that is composed of 3-cycles and 4-cycles. During loading, contacts are lost and longer cycles arise until only a single 9-cycle remains. (b) One way to quantify these structural changes is by calculating the global clustering coefficient $C$ (solid curve), which undergoes a sharp drop at peak stress, signifying the onset of material failure. (The dashed curve shows the standard deviation of the distribution of local clustering coefficients $C_i$). (c) A decrease in mean subgraph centrality $Y$ (solid curve) also illustrates the loss of short cycles during deformation. More specifically, the mean network bipartivity $R$ (dashed curve) increases with axial strain, highlighting that, during loading, closed walks of even (respectively, odd) length contribute more (respectively, less) to the mean subgraph centrality of the contact network. We adapted this figure, with permission, from [58].

cycle basis and then constructed cycle-participation vectors (see Section 2.2.3) from a contact network after each strain step (i.e., at each strain state) during loading. They observed that temporal changes in the cycle-participation vectors of the particles between consecutive strain steps are correlated positively with temporal changes in kinetic energy over those steps. They also observed that large values in the temporal changes of particle cycle-participation vectors and particle subgraph centrality occur in the shear-band region. Walker et al. [80] also studied a minimal cycle basis and corresponding cycle-participation vectors to examine structural transitions in a 3D experimental granular system of hydrogel spheres under uniaxial compression. As pointed out in [79], developing quantitative predictors that are based on topological information alone is extremely important for understanding how failure and rearrangements occur in systems in which energy or force measurements are not possible.

Examining cycles in contact networks can also shed light on the behaviour of force chains. The stability, load-bearing capacity, and buckling of force chains depend on neighbouring particles (so-called
spectator grains) to provide lateral support \[19, 20, 229\]. Because 3-cycles appear to be stabilizing features, it is interesting to consider the co-evolution of force chains and 3-cycles in a contact network. Such an investigation requires a precise definition of what constitutes a force chain, so that it is possible to (1) extract these structures from a given packing of particles and (2) characterize and quantify force-chain properties. Several definitions of force chains have been proposed; see, for example, \[14, 16, 27, 230\]. The studies that we describe in the next three paragraphs used a notion of ‘force chains’ from \[14, 231\], in which force chain particles are identified based on their particle-load vectors (where each particle is associated with a single particle-load vector that indicates the main direction of force transmission). More specifically, a single chain is a set of three or more particles for which the magnitude of each of their particle-load vectors is larger than the mean particle-load vector magnitude over all particles, and for which the directions of the particle load vectors are, within some tolerance, aligned with one another (i.e., they are ‘quasilinear’). We note that it is important in future work to conduct network-based studies of force-chain structure for different definitions of force chains and investigate if there are differences in their associated network properties.

Using DEM simulations of a densely packed system of polydisperse disks under biaxial loading—i.e., compressed quasistatically at a constant strain rate in the vertical direction, while allowed to expand under constant confining pressure in the horizontal direction—Tordesillas et al. \[84\] quantified the co-evolution of force chains and 3-cycles in several ways. For example, they computed a minimal cycle basis (see Section 2.2.3) of a contact network and then examined (1) the ratio of 3-cycles to the total number of cycles in which particles from a force-chain participate and (2) the force chain’s 3-cycle concentration, which they defined as the ratio of 3-cycles involving force-chain particles to the total number of particles in the force chain. When averaged over all force chains, the above two measures decrease rapidly with increased loading. Additionally, Tordesillas et al. \[84\] observed that force chains that do not fail by buckling (see \[200\] for how ‘buckling’ was defined) have a larger ratio of 3-cycle participation to total cycle participation than force chains that do buckle. Tordesillas et al. \[218\] observed, in both DEM simulations of biaxial loading (see above) and 2D photoelastic disk experiments under pure shear, that a particular measure (developed by Bagi \[228\]) of structural stability of force chains is correlated positively with the mean of the local clustering coefficient [see Eq. (2.12)] over force-chain particles. They also pointed out that 3-cycles are more stable structures than cycles of longer length during loading and that force chains with larger 3-cycle participation tend to be more structurally stable. These observations suggest that cycles—and especially 3-cycles—in contact networks are stabilizing structures that can provide lateral support to force chains. It would be interesting to study these ideas further, and to relate them to structural rigidity theory (see Fig. 10 and Section 3.1.2), especially in light of the difference between 3-cycles (which are rigid) and deformable cycles (e.g., 4-cycles).

Cycles have also been examined in granular contact networks that were extracted from DEM simulations of 3D, ellipsoidal-particle systems subject to triaxial compression \[219\]. Similar to the aforementioned results from simulations of 2D systems with disk-shaped particles, the number of 3-cycles in a minimal cycle basis of the contact networks (and the global clustering coefficient [see Eq. (2.13)]) initially decrease and then saturate with increasing load, and particles in force chains have a larger number of 3-cycles per particle than particles that are not in force chains. Tordesillas et al. \[219\] also observed that the set of 3-cycles that survive throughout loading tend to lie outside the strain-localization region (where force-chains buckle). The dearth of 3-cycles in certain regions in a material may thus be a signature of strain-localization zones. Another article to note is \[210\], which examined and compared the temporal evolution of cycles (and several other contact-network diagnostics) in a set of DEM simulations using a variety of different material properties and boundary conditions.
In another interesting study, Walker et al. [220] examined the phenomenon of aging [232, 233]—a process in which the shear strength and stiffness of a granular material both increase with time—in collections of photoelastic disks subject to multiple cycles of pure shear under constant volume. Because aging is a slow process, it can be difficult both to uncover meaningful temporal changes in dynamics and to characterize important features in packing structure that accompany aging. To overcome these challenges, Walker et al. [220] first analysed the time series of the stress ratio (using techniques from dynamical-systems theory) to uncover distinct temporal changes in the dynamics of the system. (See [220] for details.) After each small, quasistatic strain step, they also extracted the contact network of the packing at that time to relate aging to changes in topological features of the network structure. As the shear-jammed regime is approached during prolonged cyclic shear, they observed, on average, that force chains are associated with more 3-cycles and 4-cycles from the minimal cycle basis.

We have just discussed many articles that concern transitions in granular matter from a solid-like regime to a liquid-like regime. One can also use changes in the loop structure of a contact network to describe the opposite transition, in which a granular material transitions from an unconstrained, flowing state to a solid-like state during the process known as jamming (see Section 3.1.1). Studying 2D frictional simulations of isotropically compressed granular packings, Arévalo et al. [73] examined a granular contact network as they varied packing fraction. They observed that the number of cycles (which were called polygons in [73]) in the contact network grows suddenly when the packing fraction approaches the critical value \( \phi_c \) that marks the transition to a rigid state (see Fig. 12). They also observed that 3-cycles appear to be special: they continue to grow in number above the jamming point, whereas longer cycles slowly decrease in number after jamming. Although they observed a non-linear relationship near the jamming point between \( Z \) (the contact number, which is the usual order parameter for the jamming transition) and the number of 3-cycles [83], these quantities appear to depend linearly on each other after the transition. These results suggest that one can use the evolution of contact loops to understand the transition to a rigid state and to characterize subsequent changes in the system.
Application to tapped granular materials.

Properties of contact networks have also been used to study tapped granular materials, in which a packing of grains is subject to external pulses of excitation. In most studies of tapped granular materials, the packing and pulses are both oriented vertically. The intensity $\Gamma$ of these mechanical perturbations (so-called ‘taps’) is usually quantified as a dimensionless ratio of accelerations, such as the ratio of the peak acceleration of the excitation to the acceleration of gravity [236, 237]. Tapped granular materials are interesting because the packing fraction $\phi$ is not necessarily a monotonic function of the tapping intensity $\Gamma$ [238–240]. Instead, it reaches a minimum value $\phi_{\text{min}}$ at an intensity of $\Gamma_{\text{min}}$, and it then increases as the tap intensity increases (see Fig. 13a). Consequently, one can achieve steady states with the same packing fraction by using different tap intensities (i.e., both a ‘low’ tap intensity, which is smaller than $\Gamma_{\text{min}}$, and a ‘high’ tap intensity, which is larger than $\Gamma_{\text{min}}$). These steady states are not equivalent to each other, as they have different force-moment tensors [237], for example. An interesting question is thus the following: What features of a granular packing distinguish between states at the same packing fraction that are reached by applying different tap intensities?

Recent work has suggested that properties of contact networks—especially cycles (which, in this case, are particle contact loops)—can distinguish between steady-state configurations at the same packing fraction that are generated from different tap intensities. For example, in simulated 2D granular packings subjected to tapping [235, 241] (see Fig. 13b), as $\Gamma$ is increased in the regime $\Gamma < \Gamma_{\text{min}}$, the number of 3-cycles (i.e., triangles) and the number of 4-cycles (i.e., squares) both decrease. As $\Gamma$ is increased in the regime $\Gamma > \Gamma_{\text{min}}$, the opposite trend occurs, so the numbers of 3-cycles and 4-cycles increase. This makes it possible to differentiate configurations at the same $\phi$ obtained from low and high tap intensities. (See Fig. 13e and f for a plot of the number of triangles versus $\Gamma$ and $\phi$.) However, geometrical measures like the pair-correlation function, distributions of Voronoi tessellation areas or bond orientational order parameters do not seem to be as sensitive to differences in these two different states of the system (see Fig. 13c and d), perhaps because they quantify only local proximity rather than directly examining contacts. (See [235] and references therein for details of these descriptors.) These results suggest that topological features of a contact network can capture valuable information about the organization of granular packings.

3.1.4 Other subgraphs in contact networks. When studying contact networks, it can also be helpful to explore network motifs other than cycles. Recall from Section 2.2.6 that motifs are subgraphs that appear often in a network (e.g., in comparison to a null model) and are thus often construed as being related to system function [96, 99, 242, 243]. Network motifs, which traditionally refer to small subgraphs, are a type of mesoscale feature, and it can be insightful to examine how their prevalences change in a granular material as it deforms.

One system in which motifs and their dynamics have been studied is frictional, bidisperse, photoelastic disks subject to quasistatic cyclic shear [110]. After each small strain increment (i.e., strain step) in a shear cycle, Tordesillas et al. [110] considered the contact network of the granular packing. For each particle $i$ in the contact network, they extracted the subgraph of particles (nodes) and contacts (edges) formed by the central particle $i$ and particle $i$’s contacting neighbours. This process results in a set of $N$ subgraphs (which, borrowing terminology from [110], we call conformation subgraphs), where $N$ is the number of particles in the network.

---

9 In some degenerate cases (e.g., when there is a network component that consists of a clique), this includes duplications of subgraphs, even when the nodes of a network are labelled.
Fig. 13. Using a contact network to distinguish states of the same packing fraction in simulations of tapped granular materials. (a) Packing fraction $\phi$ versus tap intensity $\Gamma$. The horizontal lines connect states at the same packing fraction that were obtained using different tap intensities. (b) A section of one of the packings. (c) The mean value of the 'bond orientational order parameter' (or simply 'bond order parameter') as a function of tap intensity. The bond order parameter, which is often used to quantify local and long-range crystalline structure in a material [234], was computed on each subgraph of particles defined by a central particle and the set of its neighbours within a distance of 1.2 particle diameters, and it was then averaged over all such subgraphs to obtain a mean value. (d) The mean value of the bond order parameter for different values of packing fraction, where the arrows indicate the direction of increasing $\Gamma$. It is difficult to differentiate between different states at the same packing fraction using this quantity. (e, f) The same as panels (c,d), but now the vertical axis is the number of 3-cycles in the contact network. Calculating the number of 3-cycles successfully separates different states of the system with the same packing fraction. We adapted this figure, with permission, from [235].
To examine packing rearrangements as a system is sheared, Tordesillas et al. [110] represented each distinct conformation subgraph present at any time during loading as one ‘state’ in a Markov transition matrix, and they studied transitions between the conformation subgraphs as a discrete-time Markov process. More specifically, each element in the $n_c \times n_c$ transition matrix (where $n_c$ is the total number of unique conformation subgraphs and hence the number of unique states) captured the fraction of times that each conformation subgraph transformed to each other conformation subgraph (including itself) after four quasistatic steps of the shearing process. Tordesillas et al. [110] reported that force-chain particles typically occur in network regions with high mean degree, high mean local clustering coefficients, and many 3-cycles. (Note that this study, as well as the others that we describe in the present and the following paragraph, define force-chains as in [14, 231].) Furthermore, when considering the conformation subgraphs of particles in force chains that fail by buckling (see [227, 229] for details on the definition of ‘buckling’), the most likely transformations tend either to maintain the topology of those conformation subgraphs or to involve specific types of transitions. Specifically, conformation subgraphs in which the central particle has a larger degree or is part of more 3-cycles transition to conformation subgraphs in which the central particle has smaller degree or participates in fewer 3-cycles. Tordesillas et al. [110] also used force information to compute a measure of structural stability (based on a structural-mechanics framework [218, 228] and summarized in a single number) for each conformation subgraph. They then split the full range of the stability values into several smaller ‘stability intervals’ (i.e., small ranges of contiguous structural-stability values) and modeled transitions between stability intervals as a Markov chain. They examined the number of conformation subgraphs that occupy each stability interval and observed pronounced peaks in some intervals that persist during loading. They also reported that conformation subgraphs whose central particles belong to force chains tend to be more stable and that conformation subgraphs whose central particles are part of buckling force chains have a higher probability of transitioning from high-stability states to low-stability states than vice versa. (For details, see Fig. 7 of [110] and the corresponding discussions.)

Walker et al. [80] used similar methods to study self-assembly in an almost-frictionless 3D system of hydrogel spheres under quasistatic, cyclic uniaxial compression (see Fig. 14a–c). After every compression step, they constructed the contact network for the system and examined two types of subgraphs for each particle: (1) conformation subgraphs, which (as discussed earlier) consist of a single central particle $i$ and that particle’s contacts; and (2) the cycle-participation vector of each particle (see Section 2.2.3). Walker et al. [80] determined the set of all unique conformation subgraphs that existed during the above compression process. They then used each of those conformation subgraphs as one state in a transition matrix, the elements of which give the fraction of times (across the whole experiment) that a particle in one state transitions to any other state in consecutive compression steps. To focus on the presence or absence of a particle in an $l$-cycle (using cycle lengths up to $l = 10$), they binarized each element of the cycle-participation vectors. (The resulting vectors thus indicate, for each particle, whether it is part of at least one $l$-cycle.) Walker et al. [80] then constructed a transition matrix in which each state is a unique binarized cycle-participation vector that occurs during the experiment. The two transition matrices capture useful information about the most likely transformations that occur between different conformation subgraphs and cycle-participation vectors as one compresses or decompresses the granular system. For both types of mesoscale structures, Walker et al. [80] used their transition matrices to extract almost-invariant sets, which indicate sets of conformation subgraphs or cycle-participation vectors (i.e., states) that tend to transition among themselves more than to states in another almost-invariant set. (See [80] for details.) In Fig. 14d, we show the most common conformation subgraphs in each almost-invariant set of the conformation subgraphs. The conformation subgraphs formed by force-chain particles belong mostly to Set 3 (see Fig. 14d), which consists of densely-connected conformation
Fig. 14. Conformation subgraphs can help quantify structural transitions in a 3D network of hydrogels. (a) A rendering of a granular system composed of spherical, hydrogel beads [244, 245] that are subject to multiple cycles of compression along a single axis. Redder colours indicate stronger contact forces. (b) An example of a contact network in a decompressed state and (c) an example of a contact network in a compressed state. (d) Pictorial representations of the most common conformation subgraphs in each almost-invariant set. The almost-invariant sets are collections of conformation subgraphs that, during loading, tend to transition amongst themselves more than they transition to conformation subgraphs in another almost-invariant set. (See the main text and [80] for details.) (e) The evolving cardinalities of almost-invariant sets can track structural transitions during cycles of compression and decompression. (The number of elements in an almost-invariant set at a given time step is the number of particles in the system whose induced conformation subgraph belongs to that set.) In highly-compressed states, most conformation subgraphs in a packing are members of Set 3, whereas most conformation subgraphs belong to Set 4 for the least-compressed states. (f) An example of a possible transition pathway, which consists of a sequence of conformation subgraphs in a given almost-invariant set and the transitions between them that occur during loading. In this example, the pathway corresponds to the following sequence of transitions between sets: Set 3 → Set 2 → Set 4 → Set 2 → Set 3. The first conformation subgraph (40) corresponds to the most prevalent conformation subgraph in Set 3, and the conformation subgraphs for the subsequent transitions between sets are those with the largest transition probabilities. For example, of all conformation subgraphs in Set 2, subgraph 19 is the one to which conformation subgraph 40 (of Set 3) is most likely to transition. These transition pathways can potentially inform constitutive-modelling efforts. We adapted this figure, with permission, from [80].

subgraphs in which there are many contacts between particles. To characterize structural changes that occur in a packing as it moves towards or away from a jammed configuration, Walker et al. [80] tracked the number of conformation subgraphs (and cycle-participation vectors) in each almost-invariant set across time. Figure 14e illustrates the temporal evolution of the numbers of elements in the almost-invariant sets of the conformation subgraphs. Walker et al. [80] also proposed transition pathways (see Fig. 14f) that may be useful for thermo-micro-mechanical constitutive-modelling efforts [246]. (A transition pathway consists of a sequence of conformation subgraphs in different almost-invariant sets, and transitions between them.)

Another way to study various types of subgraphs in granular materials is through the classification of superfamilies [98, 107] (see Section 2.2.6). A recent investigation by Walker et al. [109] considered superfamilies that result from examining 4-particle subgraphs (see Fig. 15a) in a variety of different
Fig. 15. Examination of superfamilies in a granular material. (a) One can use a rank-ordering of relative frequencies of sets of 4-node subgraphs to define a superfamily in a contact network. (b) Mean degree (blue) and stress ratio (red) versus axial strain in a DEM simulation of a deforming granular material. (The stress ratio is the minimum stress divided by the maximum stress during a cycle of loading.) The symbols at each strain increment correspond to the superfamily of the associated contact network at that point. During loading, the system is characterized by a superfamily transition (specifically, ABDECF $\rightarrow$ ADBECF) that corresponds to a shift from one with a higher prevalence of a subgraph that includes a triangle (B) to one with a higher prevalence of a subgraph without this stabilizing feature (D). We adapted this figure, with permission, from [109].

granular systems, including experimental packings of sand and photoelastic disks, and DEM simulations for different types of loading and in different dimensions. In their study, the authors defined a superfamily as a set of networks in which the prevalence of the different 4-node subgraphs have the same rank-ordering. (They did not consider whether the subgraph was a motif in the sense of occurring more frequently than in a random-graph null model.) Despite the diversity of system types, they observed several trends in the transitions between superfamilies that occur as a system transitions from a pre-failure regime to a failure regime. The most important change in the superfamilies appears to be a switch in relative prevalence from 4-edge motifs with 3 edges arranged as a triangle to acyclic 3-edge motifs (see Fig. 15). This observation highlights the important role that small mesoscale structures can play as building blocks in granular systems. It also suggests that examining the prevalence and temporal evolution of such motifs can (1) help characterize the macroscopic states of a granular system and (2) help quantify what structural changes occur as a system transitions between different states.

Notably, although calculating the prevalence of cycles and small motifs can be useful for gaining insights into contact-network structure, it can also be helpful to employ other types of network analysis that examine structure on larger scales. For example, in simulations of 2D packings of disks under isotropic compression, Arévalo et al. [73] observed that the mean shortest-path length [see Eq. (2.10)] of a contact network reflects both changes in the organization of a packing as one approaches the jamming point and changes that occur after the jamming transition takes place. The path length appears to reach a maximum value at a packing fraction below $\phi_c$. With further increases in $\phi$ below $\phi_c$, the path length then decreases rapidly, likely due to the formation of new contacts that shorten the distance between particles as the system nears the jamming point. After the jamming transition, the path length decreases further.

Before moving on, we note that because it can be difficult to measure force information accurately in many experimental granular systems, continuing to develop relevant measures for studying contact topology (i.e., without incorporating weights) remains an important area of investigation.
3.2 Force-weighted networks

Although studying only a contact network can be informative (see Section 3.1), incorporating more information into network representations allows one to more fully capture the rich behaviour of granular materials. Many of the approaches for quantifying unweighted networks can be generalized to weighted networks (see Section 2.2), although significant complications often arise (e.g., because typically there are numerous choices for precisely how to do the generalizing). From both physics and network-science perspectives, it is sensible to construct weighted networks that incorporate information about the forces between particles. This can shed considerable light on phenomena that have eluded understanding from studying only unweighted contact networks.

One important physical motivation for incorporating information about inter-particle forces is that photoelastic-disk experiments and numerical simulations have both highlighted that, particularly just above isostaticity (see the bottom of Section 2.2.1), loads placed on granular systems are not shared evenly across all particles. Instead, forces are carried primarily by a backbone of force chains. It has often been claimed that the statistical distribution of the forces are approximately exponential [8], but in fact, it depends on whether forces are recorded at a granular system’s boundary or in its bulk [247], as well as on the loading history [17]. Illuminating how force-chain structures arise, and how their structure varies across different regions of a material or under different conditions, provides crucial information for understanding how one can control the elastic modulus and mechanical stability [173] and acoustic transmission [45] in granular materials.

However, despite the ability of humans to see force chains easily in photoelastic images, it is difficult to characterize quantitatively what is or is not a force chain, and it can also be difficult to quantify how force chains evolve under compression or shear. Part of the challenge lies in the fact that force chains are spatially anisotropic, can exhibit long-range spatial correlations [17], and can have complex temporal fluctuations [16, 38]. Consequently, understanding emergent phenomena in granular systems is typically difficult using continuum theories or approaches based only on local structure. On the other hand, a network-theoretic perspective provides a fruitful way to explore interesting material properties and organization that arise from inter-particle contact forces in granular materials. Importantly, in addition to data from simulations, multiple techniques now exist for measuring inter-particle forces between grains in experiments; these include photoelasticity [17, 248], X-ray diffraction measurements of microscopic [249] or macroscopic [250, 251] deformations, and fluorescence with light sheets [244, 245]. As we will see, incorporating information about inter-particle forces into network-based investigations has yielded fascinating insights into the organization and collective structure in granular packings in both numerically-simulated and experimental systems.

The most common method for constructing a network that captures the structure of forces in a granular system is to let a node represent a particle and then to weight the edge between two particles that are in contact according to the value of the force between them. One can describe such a network with a weighted adjacency matrix (see Section 2.1) $W$ with elements

$$W_{ij} = \begin{cases} f_{ij}, & \text{if particles } i \text{ and } j \text{ are in contact,} \\ 0, & \text{otherwise,} \end{cases} \quad (3.4)$$

where $f_{ij}$ is the inter-particle force between particles $i$ and $j$. Such a force network also encodes the information in the associated contact network, and one can recover a contact network from a force-weighted network by setting all non-zero weights in the latter to 1. Although most work has determined edge weights using the normal component of the inter-particle force, one can alternatively weight the
edges by inter-particle tangential forces. With the advent of high-performance computational capabilities, one can determine inter-particle forces from DEM simulations [252] of hundreds to millions of particles. In experiments, it is possible to determine inter-particle forces using photoelastic disks (in 2D) [248] or X-ray tomography (in 3D) [253], although typically these techniques are limited to systems of hundreds to thousands of particles. In Fig. 16, we show an example of a compressed collection of photoelastic disks and illustrate the structure of a corresponding force-weighted network for the associated packing.

We now review network-based approaches for investigating force-weighted networks constructed from granular materials and discuss the resulting insights that these approaches have provided into the physical behaviour of such systems. We label most of the following subsubsections according to the type of employed methodology, although we also include a subsubsection about some specific applications to different systems.

3.2.1 Examining weighted cycles and other structural features. In Section 3.1, we discussed why examining cycles can be useful for studying granular contact networks. It is also useful to examine cycles when investigating force networks, which are weighted. For example, Smart and Ottino [81] studied the evolution of weighted contact loops in a simulation of a quasistatically tilted granular packing. They used topological information (i.e., which particles are in contact) to define the presence of a cycle, and they defined a notion of loop stability,

\[ \xi_l = \frac{1}{\bar{f}} \prod_{i=1}^{l} f_{\text{edge}}^i, \quad (3.5) \]

to quantify the range of compressive loads that a given loop can support. In Eq. (3.5), \( l \) is the number of edges in the loop (i.e., its length), \( f_{\text{edge}}^i \) is the contact force of the \( i \)th edge, and \( \bar{f} \) is the mean edge weight (i.e., mean force) over all of the edges in the loop. See Fig. 17 for a schematic of this stability measure for a 3-cycle. For \( l = 3 \), the quantity \( \xi_3 \approx 1 \) corresponds to having approximately equal contact forces on all edges and is the most stable configuration (see Fig. 17a). The value of \( \xi_3 \) approaches 0 as the contact...
Fig. 17. A measure of the stability of 3-cycles quantifies the effects of tilting a granular packing. The loop stability $\xi_3$ of a 3-cycle is determined from the contact forces along each edge of the loop. (See the main text for details.) (a) A contact loop with approximately equal forces on all edges is very stable to perturbations in the direction of the compressive force vectors on the loop, and has a loop stability that is close to 1. (b) A contact loop in which one of the edges has a much smaller force than the others is very unstable to perturbations; it has a loop stability near 0. (c) The normalized mean loop stability $\xi^*_3$ tends to decrease with increasing tilting angle $\theta_g$. We adapted this figure, with permission, from [81].

A force on one edge becomes much smaller than those on the other two edges. As illustrated in Fig. 17b, this situation is unstable. Both the density of 3-cycles (specifically, the number of 3-cycles in the system divided by the total number of particles) and a normalized 3-cycle loop stability

$$\xi^*_3 = \frac{\langle \xi_3(\theta_g) \rangle}{\langle \xi_3(\theta_g = 0) \rangle}$$

(where the brackets denote means over all 3-cycles in a network) tend to decrease with increasing tilting angle $\theta_g$ (see Fig. 17c). Smart and Ottino [81] also reported that the effect of tilting on loop stability is largely independent from the effect of tilting on mean coordination number (i.e., mean degree).

Tordesillas et al. [84] examined what they called a force cycle, which is a cycle of physically-connected particles in which each contact carries a force above the global mean. Using DEM simulations of a biaxially compressed, dense granular system—the sample was compressed quasistatically at a constant strain rate in the vertical direction, while allowed to expand under constant confining pressure in the horizontal direction—they studied the evolution of 3-force cycles (i.e., force cycles with 3 particles) in a minimal cycle basis of the contact network with respect to axial strain. They observed that 3-force cycles initially decrease in number during strain hardening, before increasing in number at the onset of force-chain buckling [200], and then finally leveling out in number in the critical-state regime. (See the third paragraph of Section 3.1.3 for a brief description of these different regimes of deformation.)

In Fig. 18, we show a plot of the number of 3-force cycles and the shear stress versus axial strain. The 3-force cycles that arise at the onset of buckling are often part of force chains (using the definition of force chains from [14, 231]). Additionally, these 3-force cycles tend to concentrate in the region of the shear band, where they can act as stabilizing structures both by frustrating relative rotations and by providing...
strong lateral support to force chains. However, with increased loading, the system eventually fails, and Tordesillas et al. [84] suggested that the increase in the number of 3-force cycles may be an indicator of failure onset. Qualitatively similar results were observed when examining the evolution of 3-force cycles in three DEM simulations [210], each with slightly different material properties and boundary conditions, and in DEM simulations of 3D ellipsoidal-particle packings subject to triaxial compression [219].

Using DEM simulations for biaxial compression similar to those described in the previous paragraph, Walker and Tordesillas [58] examined the evolution of force-weighted networks with axial strain using several of the network concepts that we discussed in Section 2.2. Unsurprisingly, they found that including contact forces in their network analysis yields a more complete characterization of a granular system than ignoring them.

One measure that they felt was particularly useful is a weighted version of subgraph centrality (see Section 2.2.5). From a contact network, Walker and Tordesillas [58] first extracted all conformation subgraphs. As described in Section 3.1.4, a conformation subgraph is a subgraph that consists of a given particle $i$ and that particle’s immediate contacts. (Each particle in a network thus yields one conformation subgraph.) To incorporate inter-particle force information, Walker and Tordesillas [58] generated force-weighted conformation subgraphs by weighting each edge in the conformation subgraphs by the magnitude of the normal force component along that contact. They then computed a weighted subgraph centrality $Y^w_i$ for each force-weighted conformation subgraph, and they computed changes $|\Delta Y^w|$ in magnitude between consecutive strain steps of a suitably averaged version ($\tilde{Y}^w$) of this quantity (see [58] for details). They observed that the temporal evolution of $|\Delta Y^w|$ with strain step is effective at tracking large changes in energy dissipation rate that can occur due to rearrangement events (e.g., force-chain buckling) associated with the loss of inter-particle contacts. They also observed that the central particle in the conformation subgraphs that undergo the largest changes in weighted subgraph centrality seems to be associated with locations with much dissipation, such as in the shear band and in buckling force chains or the neighbouring particles of those force chains. (See [14, 231] for the employed definition of force chains and [200] for the employed specification of force-chain buckling.)
Walker and Tordesillas [58] highlighted that network analysis—and especially the examination of mesoscale features—can be helpful for gaining insights into mechanisms that regulate deformation in granular materials. Such studies can perhaps also help guide efforts in thermo-mechanical constitutive modelling [254].

3.2.2 Extracting multiscale architectures from a force network using community detection. A major benefit of studying a network representation of a granular system (and using associated computational tools) is that it provides a natural framework in which to probe structure and dynamics across several spatial scales. One can examine different spatial scales in multiple ways, including both physically (e.g., using distance in Euclidean space or in some other metric space) or topologically (e.g., using the hop distance along edges in a network). The ability to successfully study mesoscale architecture, which manifests at intermediate scales between particle-level and system-wide organization, is an especially important contribution of network analysis. One of the most common ways to examine mesoscale structures is with community detection (see Section 2.2.7), which one can use to extract sets of densely-connected nodes in a network [31, 32]. One can also tune community-detection methods to examine sets of nodes of a range of sizes, from very small sets (with a lower limit of one node per set) to large sets (with an upper limit of all nodes in a single set).

By applying multiscale community-detection methods to force-weighted contact networks of photelastic disks and numerical simulations of frictionless packings, Bassett et al. [27] identified chain-like structures that are visually reminiscent of force chains in granular packings (see Fig. 19a). Notably, the algorithmic extraction of these ‘distributed’ mesoscale structures [27, 112, 113, 255], in contrast to ‘compact’, densely-connected geographical domains in a material [21], required the development of a geographical null model, which can be used in modularity maximization and which encodes the fact that a given particle can exert force only on other particles with which it is in direct contact [126] (see Section 2.2.7). The different type of mesoscale organization extracted by this geographical null model highlights the fact that using physically motivated network-based approaches, which incorporate spatial and/or physical constraints, may give different information about a granular system than what one obtains when doing network calculations based only on network structure. In a modularity-maximization approach to community detection, one can also tune a resolution parameter of a modularity objective function to identify and characterize network communities of different sizes (e.g., as in Fig. 19a). One can also use inference procedures, in concert with community detection or other forms of clustering, to determine particularly important scales in a network.

One interesting result from [27] is that properties of force chain-like communities can distinguish frictional, laboratory packings from frictionless, simulated ones, allowing a quantification of structural differences between these systems (see Fig. 19b). In later work, Huang and Daniels [255] used similar techniques to examine friction-dependence and pressure-dependence of community structure in 3D simulations of compressed granular materials. To further quantify such mesoscale organization and examine how it changes with compression, Giusti et al. [112] extracted communities using the geographical null model, and used ideas from algebraic topology to define a topological compactness factor that quantifies the amount of branching—versus compact, densely-interconnected regions—in communities of a force network from 2D granular systems. The approach from [27] was extended to multilayer networks (see Section 2.2.7) in [113], providing a way to link particulate communities across compression steps (rather than extracting new ones at each step) when examining how such communities reconfigure. These studies helped lay groundwork to improve understanding of how the multiscale nature of force-chain architecture impacts bulk material properties. Various community-detection approaches have also been used for identifying other types of inhomogeneities in granular matter [256].
Fig. 19. Community-detection techniques can uncover multiscale force-chain structures in laboratory and simulated granular packings. (a) Examples of force-chain communities extracted using modularity maximization for networks constructed from 2D laboratory and simulated granular packings. Tuning a resolution parameter allows one to detect mesoscale features at multiple scales in a packing: smaller resolution-parameter values yield larger and more compact communities, and larger values yield communities that are smaller and more chain-like. (b) One can characterize community structure using different measures, which are able to differentiate between laboratory and simulated packings in a quantitative manner. In this figure, the size of a community is the number of particles in that community, the network force of a community is the contribution of that community to modularity, and the gap factor is a diagnostic that measures the presence of gaps and the extent of branching in a community. (The gap factor is small for very linear or compact communities, and it is large for communities with curves and/or branching.) We adapted this figure, with permission, from [27].

Before moving on, it is important to note that—although related—the definition of force chain structure using the community-detection approaches that we described above [21, 27, 112, 113, 255] differs from the definitions of force chains that have been used in some other studies (e.g., see [14]). In future work, it is important to examine how the properties of force chains differ when they are defined in different ways.

### 3.2.3 Some applications.

**Comminution processes.** A network-based approach can give fascinating insights into comminution, the fragmentation of a material into smaller pieces. Walker *et al.* [257] used DEM simulations to study comminution in a granular material under uniaxial compression and reported that the degree distribution of the system’s contact network (which we recall is unweighted) evolves towards a power law during this process. This observation is consistent with the development of a power-law grain-size distribution, in which large particles are hubs that have many smaller, neighbouring particles, which make up the majority of a packing. Walker *et al.* [257] also examined several other features (such as measures of network efficiency, node betweenness, and cycle populations) of both contact networks and networks weighted by the normal force between particles for progressively larger strains to examine what changes occur in a granular system during comminution.

**Heat transfer.** Another problem that has been examined using network-based methods is heat transfer in granular matter. Using a heat-transport model on simulations of a compressed granular material, Smart *et al.* [24] probed the effects of heterogeneity in the force distribution and the spatial arrangements of forces in a system on heat transfer through the material. Specifically, they compared measures of
transport in the (normal) force-weighted network of a granular system to two null-model networks with the same contact topology but with either (1) homogenous, uniform edge weights that were equal to the mean force of the packing; or (2) the same set of heterogeneous edge weights from the actual granular network, but assigned uniformly at random to the edges. Smart et al. [24] estimated the thermal diffusivity and effective conductivity from simulations on each network, and they observed that the real granular system has significantly higher diffusivity and effective conductivity than the homogenous null model. Additionally, comparing the results from the real material to the null model with randomly reassigned edge weights demonstrated that the qualitative differences between the real granular network and the homogenous null model could not be explained by the heterogeneity in the force distribution alone, as the authors observed that this second null model (with randomly reassigned edge weights) was also not a good medium for heat transfer.

To investigate what features of a granular network facilitate efficient heat transfer, Smart et al. [24] defined a weighted network distance (see Section 2.2.2) between particles \( i \) and \( j \) as \( d_{ij}^w = 1/H_{ij} \), where \( H_{ij} \) is the local heat-transfer coefficient, such that the network distance between two particles in contact is proportional to that contact’s resistance to heat transfer. Note that \( H_{ij} \propto f_{ij}^\nu \), where \( f_{ij} \) is the magnitude of the normal force between \( i \) and \( j \), and \( \nu \geq 0 \) is a constant. They then defined a network-based (but physically-motivated) measure of heat transport efficiency as the weighted efficiency \( E^w \) (see Section 2.2.2) computed using the distances \( d_{ij}^w \). In a comparison between the real granular system and the two null models, \( E^w \) gave the same quantitative results as the effective conductivity. In particular, the calculations in [24] revealed that the real granular system has a larger efficiency than that of either null model, suggesting that the spatial distribution of force-chain structure in the granular network appears to facilitate heat transport. Finally, iterative edge removals in decreasing order of geodesic edge betweenness centrality [Eq. (2.17)] yield a faster decrease in effective conductivity than either uniformly-random edge removals or edge removals in decreasing order of the local heat-transport coefficient, further illustrating the utility of network-theoretic measures for examining transport phenomena in granular systems.

**Acoustic transmission.** One can also examine the effect of network structure on properties such as electrical conductivity in systems composed of metallic particles or on other types of transport (such as sound propagation) through a particulate material. The transmission of acoustic signals through granular materials is poorly understood [23], and it is particularly challenging to explain using continuum or particulate models [42–44]. Bassett et al. [21] represented compressed, 2D packings of bidisperse, photoelastic disks as force-weighted contact networks, and found that some network diagnostics are able to identify injection versus scattering phases of acoustic signals transmitted through a granular material. Among the diagnostics that they computed, the authors observed that network efficiency (see Eq. (2.11)) in Section 2.2.2) is correlated positively with acoustic transmission during the signal-injection phase, suggesting that high-amplitude and high-energy signals are transmitted preferentially along short paths (and perhaps even shortest paths) through a force-weighted contact network. In contrast, low-amplitude and low-energy signals that reverberate through a packing during the subsequent signal-scattering phase correlate positively with the intra-community strength \( z \)-score, which characterizes how strongly a node connects to other nodes in its own community. These results suggest that one can use network diagnostics that probe diverse spatial scales in a system to describe different bulk properties. Because [21] did not use community-detection approaches informed by a geographical null model (see Section 2.2.7), it did not address (and it is not yet fully understood) how acoustic transmission depends on the multiscale architecture of chain-like structures reminiscent of force chains. This remains an open issue, and network-based approaches—for example, using geographical null models and other ideas that pay attention to the role of space—are likely to be important in future work on this topic.
Fig. 20. The contacts and forces in a granular network at two different force thresholds. (a) When only contacts with forces $f > 1.3 \langle f \rangle$ (where $\langle f \rangle$ is the mean force) are included, the resulting network is composed of many disconnected components. (b) As one decreases the threshold, the components grow, and if only contacts with $f > \langle f \rangle$ are included, the network is connected and ‘percolates’ along the axis of compression. The system is a numerical simulation of a dense, 2D collection of hard spheres under biaxial compression. We adapted this figure, with permission, from [19].

3.2.4 Thresholded force networks. Before researchers started using network-based methods as a common perspective for studying granular materials, Radjai et al. [19, 258] reported that under stress, the force network in simulations of granular matter organizes into two subsets: one set with ‘strong’ contacts and another set with ‘weak’ contacts. The strong subnetwork of forces forms a backbone of chain-like structures that carry most of a system’s load and tend to align approximately with the direction of compression. Between these strong force chains, there is a weak subnetwork of contacts that carry forces that are less than the mean. This weak subnetwork tends to have an anisotropy that is orthogonal to the compression, and it may provide support to the backbone of strong forces. Such heterogeneity in a force network is an interesting feature of granular materials, and network-based approaches provide a direct way to examine how strong and weak contacts impact material properties and stability.

These ideas have been explored using force-thresholded networks [73, 82, 83, 259], in which one retains only contacts that carry a force of at least some threshold $f_{th}$. That is,

$$A_{ij}^{th} = \begin{cases} 1, & \text{if particles } i \text{ and } j \text{ are in contact and } f_{ij} \geq f_{th}, \\ 0, & \text{otherwise}. \end{cases}$$

(3.6)

The threshold $f_{th}$ should be lower-bounded by the smallest non-zero contact force $f_{min}$ in the system being considered. (Note that when $f_{th} = f_{min}$, one includes all contacts in the force-thresholded network.) It is common to use a system’s mean force $\langle f \rangle$ as a reference point and to systematically vary $f_{th}$ to be different fractions of the mean force. Whether one uses the normal or the tangential component of the force, varying $f_{th}$ results in a series of thresholded networks that one can subsequently characterize using traditional network or percolation-based analyses (which we discuss in the following three paragraphs of this subsubsection) or using methods from computational algebraic topology (which we discuss in Section 3.2.5). See Fig. 20 for an example depicting the contacts and forces at two different values of the force threshold in a simulation of a 2D granular material. There is also a connection between the
idea of force-thresholded networks and carrying out modularity maximization with the geographical null model [Eq. (2.23)] and a resolution parameter $\gamma$ (see Section 2.2.7). Similar to how increasing the threshold $f_{th}$ selects the subset of particles in a network for which inter-particle forces exceed or equal the threshold, in modularity maximization, increasing the resolution-parameter value $\gamma$ tends to yield communities of particles such that within a community, the inter-particle forces are at least as large as the threshold $\gamma \langle f \rangle$.

Arévalo et al. [73, 82, 83] examined several network diagnostics—for example, mean degree, shortest-path length, diameter, LCC size, and component-size distributions—with respect to the force threshold $f_{th}$ (and at different values of the packing fraction) in DEM simulations of 2D granular packings under isotropic compression. The computations in [73, 82] suggest that the way in which many of these measures change as one varies $f_{th}$ depends on packing fraction, and many of the measures can thus potentially be used to help understand differences in the organization of force networks above versus below the jamming point. For example, for packing fractions above the jamming point, the LCC size and the shortest-path length undergo qualitative changes in behaviour near a threshold $f_{th} \approx \langle f \rangle$, signifying interesting structural changes in the organization of the force networks at that threshold. The relationship between the number of 3-particle contact cycles (i.e., triangles) and the force threshold was also examined in Arévalo et al. [73, 83]. In the jammed state, they observed a steep decline in the number of 3-cycles in the networks as they increased the threshold and considered only progressively larger forces. This observation suggests that triangles (or at least one of their contacts) belong primarily to the weak subnetwork of forces that help support strong, filamentary force-chain structures. See Fig. 1 and Sections 3.1.3 and 3.2.1 for other discussions of the roles of cycles and their relationship to force chains. As we discuss in Section 3.2.5, one can also use methods from computational algebraic topology to examine the evolution of cycle organization in dense granular materials.

Another way to study force-thresholded granular networks is to use a percolation-like approach (see Section 2.2.9). For example, one can examine the sizes and number of connected components in a thresholded network with respect to $f_{th}$ [176, 180–182, 260]. For dense packings, the intuition is that when $f_{th}$ is very large, a force-thresholded granular network splits up into several disconnected components; and as one decreases $f_{th}$, these components begin to merge until eventually all contacts are in a single component. In this type of bond percolation, which edges are included in the network thus depends on the force that the edges carry. One can look for a critical threshold $f_{th}^{c}$ such that for $f_{th} > f_{th}^{c}$, the network fragments into many small components, but as one lowers the threshold towards $f_{th}^{c}$, a large, percolating cluster forms in the system (see Section 2.2.9). Quantities that are often investigated when studying this type of force-percolation transition include $f_{th}^{c}$ and 'critical exponents' for the transition (see Section 2.2.9). Because both experimental and computational granular systems are finite in practice, such investigations often use finite-size scaling techniques.

Several simulation-based studies of granular systems have deployed percolation analyses based on force-thresholded networks to quantify the organization of granular force networks, how such organization changes with increasing compression, and other phenomena [180–182, 260]. For example, Pastor-Satorras and Miguel [181] studied force percolation in the $q$-model [10] of anisotropic granular force networks, in which there is a preferred direction of force propagation. They concluded that the asymmetry in the model has a significant effect on the percolation transition, and they found that the critical exponents differ from those of isotropically compressed granular force networks. Kovalcinova et al. [180] investigated force percolation in a variety of simulations of slowly compressed, 2D granular systems. They examined the effects of polydispersity and friction, finding that these factors can qualitatively influence various features of the percolation transition. Very recently, Pathak et al. [182] also investigated the force-percolation transition in simulations of jammed granular packings at fixed pressures.
3.2.5 Methods from computational algebraic topology. In addition to traditional approaches for network analysis, one can also study the architecture of granular networks using ideas from algebraic topology. Persistent homology (PH) [153, 165, 183, 190] (see Section 2.2.10) seems especially appropriate, and over the past several years, it has provided a fruitful perspective on the structure of compressed [163–166, 261] and tapped [167–169] granular materials. Very recently, methods from computational algebraic topology have also been used to study responses of granular materials to impact [262, 263]. One way to characterize the organization and evolution of granular force networks is to examine how Betti numbers (see Section 2.2.10) change with respect to a force threshold (and also with respect to packing fraction) in compressed granular systems. This includes studying the birth and death of components (determined by $\beta_0$) and loops (determined by $\beta_1$) as one varies a force threshold by computing and analysing persistence diagrams (PDs). See Fig. 8 of Section 2.2.10. Examining when and how long different features persist in a network provides a detailed characterization of the structure of granular force networks, and one can quantify differences between two networks by defining measures of ‘distance’ between their associated PDs. These capabilities allow a PH framework to provide a distinct set of contributions that complement more traditional network-based analyses of components, cycle structure, and other features.

Kondic et al. [163] investigated how simulated 2D granular force networks evolve under slow compression as they cross the jamming point. They first demonstrated that one can identify the jamming transition by a significant change in behaviour of $\beta_0$ (specifically, there is an increase in the number of components at a force threshold approximately equal to the mean force $\langle f \rangle$), and that structural properties of the network—such as the size of the connected components—continue to change above jamming. Kondic et al. [163] also demonstrated that $\beta_0$ and $\beta_1$ can quantitatively describe the effects of friction and polydispersity on the organization of force networks (and can distinguish how friction and polydispersity alter the structure of a force network). This work was extended in Kramár et al. [164], who studied numerical simulations of 2D, slowly compressed, dense granular materials using PH. In addition to examining the values of the Betti numbers, they also computed $\beta_0$ and $\beta_1$ PDs (PD0 and PD1, respectively) as the system was compressed to quantify the appearance, disappearance, and lifetimes of components and loops. In [164], Kramár et al. defined a filtration over the clique complex of the networks (see Section 2.2.10), so only loops with four or more particles were counted. To extract useful information from the PDs, they binned the persistence points in each diagram into different regions, corresponding to features that (1) are born at any force threshold but have relatively short lifetimes compared to those that persist for a large range of thresholds and that are born at either (2) strong, (3) medium, or (4) weak forces. Their persistence analysis led to several insights about the structure of a normal-force network as a granular system is compressed, as well as insights into differences in the structure of a normal-force network for systems with different amounts of friction and different polydispersities. For example, Kramár et al. [164] observed that, near the jamming point, frictionless packings appear to have more ‘extreme’ features than frictional packings, in the sense that frictionless packings have many more $\beta_0$ persistence points that are born at either weak or strong forces and that are relatively long-lived. Kramár et al. also pointed out that the effects of polydispersity and friction may be difficult to observe using traditional measures such as the probability density function of the normal forces.

Kramár et al. [166] also used PH to study force networks from simulations of slowly compressed, polydisperse packings of disks in 2D as they traverse the jamming transition (see Fig. 21a). As they compressed the system through a range of packing fractions $\rho$, they extracted force information at approximately fixed time intervals during the simulation, and they then computed PDs of components and loops (i.e., PD0 and PD1, respectively) for each force network sampled during compression (see Fig. 21b). As in other studies, [166] used the clique complex of the force networks to avoid counting 3-particle loops. They then used the bottleneck distance $d_B$ and two variants, $d_{W_1}$ and $d_{W_2}$, of the Wasserstein
FIG. 21. Examination of granular force networks using computational algebraic topology. (a) An example of a force network from a simulation of a dense, 2D, frictional, polydisperse granular material under compression. The grayscale bar indicates the magnitude of the normal force (normalized by the mean force in the packing) between contacting particles. (b) The left panel shows PD₀ for the force network, and the right panel shows PD₁ for the force network. These PDs indicate the appearance and disappearance of connected components and loops, respectively. Because the filtration is defined over the clique complex of the force network, one counts loops only when they include four or more particles. (c) One can examine the evolution of the force network as one varies packing fraction ρ by using various distances (left axis) to quantify differences between the PDs computed from force information from consecutive samples (see the main text for details) of the system during compression. These distances (the bottleneck distance dB and two variants, dW₁ and dW₂, of the Wasserstein distance [190]) capture both local and global changes in force geometry as one compresses the system. The curve for the mean coordination number Z (right axis) gives the mean number of contacts per particle, and the steep rise in Z signifies the onset of jamming. The values of the distances (which we show for consecutive PD₀s) change dramatically as the system goes through the jamming point, after which they vary much more smoothly. See [166] for details about these measures and for an example computation of the distances between consecutive PD₁s with respect to packing fraction. (d) The ratios of some of the distance measures during compression. Considered in conjunction with the behaviour of the distance measures in panel (c), the dramatic drop in these ratios near the jamming point signifies a rapid global restructuring of the force network that occurs via many small changes in the organization of the network’s connected components. We adapted this figure, with permission, from [166].

distance [190] to quantify differences between two PDs and thereby help quantify differences in local and global features between two granular force networks. As described in Kramár et al. [166], the bottleneck distance captures only the largest difference between two PDs, whereas the Wasserstein distances include all differences between two PDs, with dW₁ more sensitive than dW₂ to small changes. (See [166] for more
details.) Calculating the various types of distance between the two $\beta_0$ PDs and the two $\beta_1$ PDs for consecutive samples (i.e., consecutive states) of a force network allows one to characterize different kinds of variations in the geometry of force networks as a function of packing fraction (see Fig. 21c). Using the aforementioned distance measures, Kramár et al. [166] observed that in the unjammed state, there can be significant (but localized) reorganization in force geometry as a packing is compressed. They also concluded that the jamming transition is characterized by rapid and dramatic global rearrangements of a force network, followed by smoother and less dramatic reconfiguration in the system above jamming, where the distances between consecutive states of a packing are much smaller than in the unjammed state. Kramár et al. [166] also found that tangential-force networks seem to exhibit similar behaviour (in most respects) to that of normal-force networks. They also observed that friction can have a significant impact on how the geometry of the forces reconfigures with compression. For example, they showed that the rate of change of loop features (as measured by the distances between consecutive PD$_s$) is larger for a frictional system than for a frictionless one below the jamming point, but just before jamming and thereafter (i.e., during the entire jammed state) differences in loop structure between consecutive packing fractions are larger for frictionless systems than for frictional ones.

In very recent work, Kondic et al. [261] used PH to examine the temporal scales on which granular force networks evolve during slow compression. They simulated dense 2D granular materials and studied the influence of the externally-imposed time scale—set by the rate of compression—on how frequently one should sample a system during compression to be able to evaluate its dynamics near the jamming transition. By varying the sampling rate and carrying out a persistence analysis to quantify the distance between consecutive sampled states of the system, their results indicate that close to jamming, a force network evolves on a time scale that is much faster than the one imposed by the external compression. See [261] for further details.

One can also use PH to study force networks in tapped granular systems. Pugnaloni et al. [169] examined DEM simulations of two different 2D systems exposed to tapping. One type of packing consisted of monosized disk-shaped particles, and the other type consisted of monosized pentagon-shaped particles. Pugnaloni et al. [169]’s investigation suggested that particle shape can play an important role in mechanical responses, which is consistent with observations from classical investigations of granular materials [18]. More specifically, Pugnaloni et al. [169] computed $\beta_0$ and $\beta_1$ with respect to force threshold in both normal-force networks and tangential-force networks. They observed for both types of force-weighted networks (but particularly for the tangential one) that the first two Betti numbers are able to clearly distinguish between disks and pentagons, where $\beta_0$ (respectively, $\beta_1$) is consistently larger (respectively, smaller) for pentagons across a wide range of force thresholds. However, using only $\beta_0$ and $\beta_1$, Pugnaloni et al. [169] were unable to clearly differentiate states with similar packing fractions but that result from different tap intensities.

In a follow-up investigation, Kondic et al. [168] simulated a series of several taps to granular packings and used PH to examine how normal and tangential force-weighted networks vary between individual tap realizations. Specifically, they computed distances between PD$_0$s and between PD$_1$s of force networks associated either with individual realizations of tapping to the same system or with individual realizations of tapping to two different systems. In one part of their study, they examined systems of disks exposed to a series of taps at two different tap intensities. (See Section 3.1.3 for a rough delineation of ‘low’ tapping intensity versus ‘high’ tapping intensity.) They observed that in terms of loop structure, the set of networks generated from a series of taps at low intensity differ far more substantially from each other—as quantified by the distribution of distances between the PD$_1$s for different realizations of the tapping—than do the set of force networks from a series of taps at high intensity. They also observed that the distances between different realizations of low-intensity tapping are as large as the distances between low-intensity tapping
and high-intensity tapping realizations. Therefore, although the high-intensity tapping and low-intensity tapping regimes yield networks with approximately the same packing fraction (see Section 3.1.3), one can use methods from PH to help explain some of the differences between the packing structure in the two regimes. In another part of their study, Kondic et al. [168] carried out a persistence analysis of tapped packings of disks and tapped packings of pentagons, and they observed clear distinctions between the two systems based on calculations of $\beta_1$ PDs. For example, for each system, they computed the PDs for a set of networks associated with several individual realizations of the same tapping intensity, and they then computed a distance between each pair of PDs for realizations within the same packing and across the two types of packings. They observed that the distribution of distances between the PDs of individual tapping realizations to the packing of pentagons is narrower and centred at a smaller value than the distribution of distances between individual realizations of taps for the packing of disks. They also observed that the distances between the disk and pentagon systems are much larger than those between different realizations of the disk system. Thus, Kondic et al. [168] were able to distinguish clearly between tapped disk packings and tapped pentagon packings using PH, especially when considering properties of loop structures. Past work using 2D experiments has also been able to distinguish between the dynamics of disk and pentagon packings using conventional approaches [18].

One can also use methods from computational algebraic topology to study granular networks in which one uses edge weights from something other than a force. For example, Ardanza-Trevijano et al. [167] used only particle positions (in the form of point clouds) and computed Betti numbers to distinguish states at the same density but that are at different mechanical equilibria. Using both experimental and simulated 2D granular packings of monodisperse particles, they constructed networks by locating the centre of each particle and then introducing a filtration parameter $\delta$, such that any two particles separated by a Euclidean distance less than or equal to $\delta$ are adjacent to each other in a graph. They considered $\delta \in [d, 1.12d]$, where $d$ is the particle diameter and the domain for $\delta$ resembles the choices that are used for determining if particles are in physical contact with each other. The authors computed, as a function of $\delta$, the first Betti number $\beta_1$ on the whole network to count the total number of loops at a given $\delta$. They also computed $\beta_1$ on the flag complex (see Section 2.2.10), thus counting the number of loops with four or more nodes at a given value of $\delta$. For values of $\delta$ that are slightly larger than $d$, Ardanza-Trevijano et al. [167] were able to separate states at the same packing fraction that are generated by tapping at different intensities. They observed for a fixed packing fraction that states that arise from lower-intensity tapping have a larger value of $\beta_1$ when computed on the whole network and a lower value of $\beta_1$ when computed on the flag complex. Their results were robust to both noise and errors in particle-position data.

3.3 Other network representations and approaches

3.3.1 Network-flow models of force transmission. Another technique for gaining insight into the organization of forces in deforming granular systems, and how microscale aspects of a force network lead to macroscale phenomena such as shear bands and material failure, is to view force transmission from the perspective of maximum-flow–minimum-cut and maximum-flow–minimum-cost problems (see Section 2.2.8). To examine a granular system using such a perspective, one can consider the ‘flow’ of force through a contact network (with some contacts able to transmit more force than others), which in turn yields a ‘cost’ to the system in terms of energy dissipation at the transmitting contacts [264–268]. One can calculate flow and costs in routes through a network (and hence determine bottlenecks in force transmission) to gain understanding of how contact structure relates to and constrains a system’s ability to transmit forces in a material. For example, Lin and Tordesillas [267, 268] constructed flow networks from DEM simulations of a system of polydisperse particles compressed quasistatically under a constant
strain rate in the vertical direction and allowed to expand under constant confining pressure in the horizontal direction. At a given axial-strain value (i.e., ‘strain state’), they assigned uniform capacities $u_{ij}$ to each edge of the contact network to reflect the maximum flow that can be transmitted through each contact, and they assigned costs $c_{ij}$ to each edge to model dissipation of energy at each contact. After each axial-strain increment during loading, they then solved the maximum-flow–minimum-cost problem for the network at the given strain state, finding that edges in the minimum cut (yielding bottlenecks in the force-transmission networks) localize in the material’s shear band. By using costs $c_{ij}$ that reflected the type of inter-particle contact (specifically, elastic contacts versus various types of plastic contacts) Lin and Tordesillas [267, 268] were able to track different stages of deformation (i.e., strain-hardening, strain-softening, and the critical-state regime). They also computed a minimal cycle basis and observed that a large majority of force-chain particles and particles in 3-cycles are involved in the set of contacts that comprise the maximum-flow–minimum-cost routes.

One can use the above approach with various definitions of force capacity and cost functions. Using simulations of the same type of system as that in the previous paragraph, Tordesillas et al. [266] constructed networks—one for each strain state as a system was loaded until it failed—that incorporated information about both the inter-particle contacts at a given strain state and the particle displacements that occur between consecutive strain steps. Specifically, if nodes $i$ and $j$ are in contact at a given strain state, the weight of the edge between them is the inverse of the absolute value of the magnitude of the relative displacement vector of particles $i$ and $j$, where one computes the displacement for each particle from the previous strain state to the current one. The distance between nodes $i$ and $j$ is 0 if the associated particles are not in contact. The intuition behind this capacity function is that, when there is more relative motion between a pair of particles, one expects those particles to have less capacity to transmit force to each other. Tordesillas et al. [265] used capacities that incorporate 3-cycle memberships of edges in a study of minimum cuts of a flow network in two samples of 3D sand under triaxial compression and in a 3D DEM simulation of simple shear. They observed that grains in the bottlenecks localize early during loading and are indicative of subsequent shear-band formation. Other work [264] studied DEM simulations of compressed, 3D bonded granular materials (where bonded signifies that the grains are connected via solid bonds of some strength) and a system of 2D photoelastic disks under shear stress with the goal of testing the hypothesis that an appropriate maximum-flow–minimum-cost approach can identify experimentally-determined load-bearing particles and force-chain particles without relying on knowledge of contact forces. Tordesillas et al. [264] examined different combinations of force-transmission capacity and cost functions, and they examined the fraction of force-chain particles that are part of the associated maximum-flow–minimum-cost network for a given capacity and cost function. In both cases, costs based on 3-cycle membership of edges seem to yield large values of these fractions, and Tordesillas et al. [264] were able to successfully forecast most of the particles that eventually become part of force chains without using information about contact forces.

\subsection{Broken-link networks.}

In previous discussions (see Sections 3.1 and 3.2), we have seen that one way to investigate the evolution of a granular system under an applied load is (1) to compute contact networks or force-weighted networks for the system with respect to packing fraction, strain, or some other control parameter and then (2) to study how different features and properties of the networks emerge and change as one varies that parameter. One can also use other network constructions to explore different mesoscale features and examine system dynamics. For example, Herrera et al. [28] designed a broken-link network (see Fig. 22) to study the dynamics of 3D granular flows. They conducted an experiment on a collection of acrylic beads immersed in a box of liquid medium, shearing the system at a constant
Fig. 22. A broken-link network can yield insights into dynamic reconfiguration in granular shear flows. (a) A schematic of the construction of a broken-link network based on changes in a proximity network that occur as a system is sheared. (b) An example of a broken-link network (with edges in green) at one instant in time. We adapted this figure, with permission, from [28].

rate \( \Omega \approx 1.05 \times 10^{-3} \) rad/s) by a rotating circular disk at the bottom of the box. (See [28] for details about their experiments.) First, they constructed proximity networks (a variant of a contact network) as a function of time. (Specifically, they constructed one network every 3-degree increment of rotation.) In their proximity network, they assigned a 'contact' (edge) to each pair of particles whose distance from one another in the given frame was within a specified distance threshold, which they chose to be a conservative upper bound for particle contact. They then defined a broken link (relative to some reference frame) as an existing edge between two particles in the reference frame that was subsequently absent in at least two later time frames (due to the particles having moved apart). A broken-link network for the frame in which a pair of particles moved apart had an edge between the two particles that were initially in contact, and broken links were not allowed to reform later. In Fig. 22a, we illustrate this procedure for constructing a broken-link network.

Studying the temporal evolution of a broken-link network provides a quantitative approach for examining particle rearrangement events in granular matter. To examine the temporal evolution of a granular system, Herrera et al. [28] examined the size of the LCC in a sequence of broken-link networks as they varied applied shear, drawing an analogy between the fraction \( \chi_b \) of broken links and the occupation probability in traditional percolation problems (see Section 2.2.9). They observed that the fraction \( s_g \) of nodes in the LCC of the broken-link network grows with \( \chi_b \) in a way that suggests that there is a continuous phase transition in \( s_g \), and they approximated the value of \( \chi_b \) at which this transition occurs. (However, as we noted in Section 2.2.9, because these networks have finite sizes, one needs to be cautious regarding statements about percolation and phase transitions.) From a physical standpoint, this transition region, corresponds to a characteristic deformation at which broken links—which are due to particle rearrangements—start to become globally connected, relative to a reference proximity network. By examining \( \chi_b \) as they applied shear to the system, Herrera et al. [28] approximated a characteristic amount of strain associated with this transition region (by mapping the value of \( \chi_b \) associated with the transition region to a corresponding value of strain), and they suggested that the determined strain scale may be useful for identifying the onset of global reorganization in the system.

In a later study, Slotterback et al. [162] used a similar approach to examine progression from reversible to irreversible dynamics in granular suspensions under oscillatory shear strain. In experiments similar to the one described above [28], the authors considered a series of 20 shear cycles of the following form: for each cycle in one experiment, a suspension was sheared with a rotating disk up to a strain amplitude of \( \theta_r \), after which the direction of rotation was reversed and the disk was rotated by \(-\theta_r\) back to its original location. The authors performed experiments for \( \theta_r \) values of 2°, 4°, 10°, 20°, and 40°. In a qualitative
sense, measuring ‘reversibility’ of the dynamics in this setup entails determining the extent to which particles return to their original positions after some number of shear cycles. To quantify this idea, one can compute the mean square displacement (MSD) of the particles after each cycle, where smaller MSD values correspond to more reversible dynamics. To study this system, Slotterback et al. [162] adjusted the idea of a broken-link network to include ‘healing’, so that broken links that are repaired in later frames do not contribute edges in the broken-link network, and they studied the temporal evolution of this broken-link network as they varied the cyclic shear and for different amplitudes $\theta_r$. In addition to the extent of spatial (ir)reversibility measured by calculating the MSD, Slotterback et al. [162] also proposed a notion of topological (ir)reversibility by examining the temporal evolution of the size of the LCC in their broken-link networks. For low values of $\theta_r$ (specifically, for $\theta_r \leq 20^\circ$), the system appears to be almost reversible: proximity-based contacts break, but they reform after shear reversal, and the fraction of particles in the LCC of the broken-link network thus grows before subsequently shrinking to almost 0 after reversal. However, for a higher shearing amplitude (specifically, for $\theta_r = 40^\circ$), the system shows signatures of irreversibility. Many broken links do not reform after a shear cycle, and after reversal, the LCC of the broken-link network remains at a value that constitutes a substantial fraction of the total system size.

3.3.3 Constructing networks from time series of node properties or from kinematic data. Another way to examine the organization of deforming granular materials is to construct networks based on the temporal evolution of particle properties [114, 116, 256, 269], an idea that draws from earlier work in complex systems on constructing networks from different kinds of time-series data. (See, for example, [107, 270–274].) In one type of construction, which yields what are sometimes called functional networks [192], one records the time series of some property of each particle and places an edge (which can potentially be weighted and/or directed) between two particles according to some relationship (e.g., some type of distance or other measure of similarity) between the particle-property time series. When using a different property, the nodes are the same, but the edges and especially edge weights will in general be different. Once networks have been constructed, one can examine them using techniques such as those in Section 2.2. Some authors have also used the generic term particle-property networks to describe networks that they constructed in some way from particle properties.

Walker and Tordesillas [114] used particle-property networks to study the evolution of DEM simulations of a quasistatically deforming, 2D granular material under biaxial compression. They constructed time series for two features (as well as their coevolution) for each particle—membership in force chains (determined as in [14, 231]) and membership in 3-cycles of a minimal cycle basis—by recording a 1 if a particle has the given property and a 0 if it does not. They then quantified the similarity of particle-property evolution using the Hamming distance [275], and they added (undirected and unweighted) edges between each particle and its $k$ closest (i.e., most similar) particles until eventually obtaining a single network with one connected component. Walker and Tordesillas [114] then extracted sets of particles that exhibit similar dynamic behaviour by detecting communities (see Section 2.2.7) in the particle-property networks. This approach uncovered distinct regions in the material—including the shear band and different subnetworks composed of primarily force chain or primarily non-force-chain particles—as well as interlaced regions in the shear band that alternate between jammed and unjammed configurations. See [116, 269] for additional studies that used membership in cycles of length up to $l = 7$ for the construction of particle-property networks.

One can also construct particle-property networks from data that do not rely on knowledge of contact or force networks. For example, Walker et al. [117] studied deformation in sand subject to plane-strain...
compression using measurements of grain-scale displacements from digital image correlations [276, 277]. From these data, they constructed *kinematic networks*, a type of network that arises from some measurement of motion (such as displacement or a rotation) over a small time increment. Walker *et al.* [117] considered each observation grid point of digital images of a sample to be a node, and they placed edges between nodes with similar displacement vectors during a small axial-strain increment. This yields a collection of (undirected and unweighted) time-ordered kinematic networks. In another study, Tordesillas *et al.* [115] calculated particle rotations and displacements for triaxial compression tests on sand using x-ray micro-tomography scanning [278]. They generated an ordered set of networks from these data for several strain steps by treating each grain as a node and linking nodes with similar kinematics during the specified interval. More specifically, they represented the displacements and rotations of each particle as points in a state space, and they connected particles that are nearby in that state space according to Euclidean distance. In the network that they constructed, each particle is adjacent to *k* nearest neighbours in state space, where *k* is as small as possible so that the (unweighted and undirected) network is using connected. Various notions of what it means to be ‘similar’, and thus how to quantitatively define edges, are discussed in [116, 117]. To probe the collective dynamics of interacting groups of particles for the network in each strain step, Tordesillas *et al.* [115] detected communities corresponding to mesoscale regions in the material that exhibit similar dynamic behaviour. Calculating the mean shortest-path length between pairs of particles in the same community yields a potentially important intermediate spatial scale (that they concluded is consistent with the shear-band diameter) of a granular system. For each strain step, they computed a variant of closeness centrality—it is similar to the one in Section 2.2.5, but it is not exactly the same—of each particle in the corresponding network, and observed that particles with large closeness centrality localize in the region of the shear band early in loading (and, in particular, before the shear band develops). This study highlights the potential of network analysis to provide early warning to detect regions of failure in particulate systems.

Methods from non-linear time-series analysis have also been used in network-based studies of stick-slip dynamics in a granular packing sheared by a slider [108]. Using so-called phase-space networks (see [107] for a description of them) to construct networks from measurements of a slider time series, Walker *et al.* [108] associated network communities with slip events.

### 3.4 Comparing and contrasting different network representations and approaches

Network representations of granular and other particulate systems, in combination with methods from network science and related disciplines, provide a plethora of ways to analyse these systems. However, for these tools to be optimally useful in furthering understanding of the physics of these systems, it is also important to draw connections between the many existing approaches. The application of network-based frameworks to the study of granular matter is relatively young, and little work thus far has focused specifically on exploring such connections, but it is crucial to determine (1) how conclusions from different approaches relate to one another and (2) how similarities and differences in these conclusions depend on the system being studied. In this short subsection, we point out a few relationships and places to begin thinking about such questions.

First, it is important to consider the network representation itself. We have discussed several different representations in this review, and some are more related to each other than others. Broadly speaking, one class of granular networks tries to encode the physical structure of a material—in the sense that edges exist only when there is an inter-particle contact—at a given point in an experiment or simulation. Such networks include contact networks (see Section 3.1) and force-weighted networks (see Section 3.2). One can obtain a contact network from an associated force-weighted network by discarding the edge weights and keeping...
information only about connectivity. Force-weighted networks thus contain much more information, and they allow one to investigate phenomena—such as force-chain organization—that may not be possible to probe with the contact network alone. However, it is still important to develop tools for the analysis of contact networks and understand what phenomena arises from features of the connectivity alone (or what one can learn and explain from connectivity alone), as force information may not always be available or may not be necessary to understand certain behaviours of a granular system. Generalizing some quantities (e.g., betweenness centralities) from unweighted networks to weighted networks also involves choices, and it is often desirable to conduct investigations that have as few of these potentially confounding factors as possible. Other types of granular networks do not encode physical connectivity, but instead directly represent something about the dynamics or changes that occur in a system during an experiment or simulation. Examples of such networks include broken-link networks (see Section 3.3.2) and particle-property networks (see Section 3.3.3). These different classes of network representations offer distinct ways of studying granular materials, and utilizing each of them should improve understanding of these systems.

For a given network representation, it is reasonable to expect that conclusions that arise from similar network quantities or methods of analysis are related to one another, whereas conclusions that result from tools designed to probe very different kinds of organization in a network provide rather different information about the underlying granular system [21, 112]. For example, some studies have suggested that in deforming granular materials, results based on calculations of clustering coefficients are similar to those from studying 3-cycles. This is intuitively reasonable, given that calculating a local clustering coefficient yields one type of local 3-cycle density. We have also observed that conclusions drawn from examinations of small subgraphs in a deforming granular system may also be related to whether or not those subgraphs contain cycles of certain lengths. As we discussed in Section 2.3, another way to draw connections between different approaches is to consider the spatial, topological, or other scales that are probed by the different approaches. For instance, in a force-weighted network, node strength is a particle-scale property, and it encompasses only very local information about a granular material. However, granular systems exhibit collective organization and dynamics on several larger scales that may be difficult to understand by exclusively computing local measures and distributions of such measures. To obtain an understanding of larger-scale structures, it is necessary to also employ different methods, such as community detection, PH, or the examination of conformation subgraphs that are composed of more than just a single particle. Utilizing such approaches has provided insights into force-chain structure, shear band formation, and reconfiguration in granular systems under load that one may not be able to obtain by considering only local network quantities. It is thus important to continue to use and develop methods to analyse granular networks across multiple scales, as doing so can provide important and new information about a system. Finally, we note that even a single approach, depending on how it is used, can provide multiple types of information about a granular network. A good illustration of this is community detection. In Section 2.2.7, for example, we saw that using different null models in modularity maximization allows one to probe different types of mesoscale architecture in granular force networks.

We look forward to forthcoming studies that directly compare the results and assumptions in different approaches (both network-based and traditional ones) and different network representations of granular and other particulate systems. Conducting principled investigations into how conclusions from various network-based approaches are related is indeed an important direction for future work.

3.5 Limitations and practicalities of simulations and experiments

Network-based studies of granular materials have examined inter-particle contact and force data (and associated dynamics, such as in the presence of external loading) from both experiments and simulations.
A recent summary of the many experimental techniques available for obtaining data about inter-particle contacts is available in the focus issue [279]. Such techniques include laser-sheet scanning [245], photoelasticity [248], x-ray tomography [253], and nuclear magnetic resonance [280]. Using each of the first three approaches, it is possible to measure both particle positions and inter-particle forces. If one is careful, it is sometimes possible to measure the forces as vectors (i.e., including both the normal and tangential components), but some techniques or systems do not have sufficient resolution to allow more than scalar or coarse-grained values. Determining the forces also helps experimentalists to confidently construct contact (i.e., unweighted) networks of particulate materials. In deciding whether or not two particles are actually in contact, rather than merely being adjacent in space, it is necessary to perform a detailed study of the effects of thresholding the data [201]. Any experimental technique will imperfectly report the presence versus absence of the weakest contacts in a system. Additionally, because of the difficulty of accessing the interior of granular materials, much more data is available for 2D force networks than for 3D force networks [279].

The most widely-used simulation techniques are DEMs [252], in which the dynamics of individual particles (usually spheres) are determined by their pairwise interactions under Newton’s laws of motion. The normal forces are typically determined from a Hertzian-like contact law (see, e.g., the sidebar in [281] for an introduction to Hertzian contacts) via an energy penalty for the overlap of two particles. The tangential (frictional) forces are most commonly modelled using the Cundall–Strack method [282] of a spring and a dashpot, but they have also been modelled via the surface roughness created by a connected set of smaller spheres [283]. For a given application, it is not known whether these simplified models capture all of the salient features of inter-grain contacts, and the situation likely differs for different applications. For example, experimental measurements of sound propagation in photoelastic disks [23] suggest that the amplitude of sound waves may be largest along a force-chain network, an effect not observed in DEM simulations [284]. This is likely a consequence of real particles physically deforming their shape to create an increased contact area through which sound can be transmitted; existing DEM simulations do not account for this effect. Another important use of particle simulations is to provide a means to investigate the robustness of network-based analyses to various amounts of experimental error [165]. Simulations provide an important check on experimental uncertainties in the determination of force-weighted networks and other network representations of granular materials. Conversely, network-based approaches provide a means to compare how faithfully simulations are able to reproduce experimental results.

4. Open problems and future directions

We now discuss a few open problems and research directions for which we anticipate important progress in the near future. We divide our comments into three main areas: the construction of different types of networks that encode various physical relationships or other properties (see Section 4.1), the application of network analysis to additional types of materials (see Section 4.2), and the application of network-based approaches to the design of materials (see Section 4.3). Network tools can provide valuable insights—both explanatory and predictive—into particulate materials and their dynamics, and a lot of fascinating research is on the horizon.

4.1 Network representations and computations

To briefly explore the potential of different approaches for constructing granular (and other particulate) networks for providing insights into the physics of granular materials (and particulate
matter more generally), we discuss choices of nodes, choices of edges, edge-to-node dual networks, multilayer networks, and annotated networks. It is also worth thinking about what calculations to do once one has constructed a network representation of a particulate system, so we also briefly consider the important issue of developing physically-informed methods and diagnostics for network analysis.

4.1.1 Definitions of nodes and edges. There are many choices—both explicit and implicit—for constructing a network [29, 287], and these choices can impact the physics that one can probe in granular networks [21]. Perhaps the most obvious choices lie in how one defines nodes and edges.

In the study of granular materials, a common definition is to treat individual particles as nodes and to treat contacts as edges (often with weights from the inter-particle forces). A natural set of open questions lies in how contact network architectures depend on different features of the grains in a system. For example, there have been several recent studies on systems composed of particles that are neither spheres nor disks—including ones with U-shaped particles [288], Z-shaped particles [289], squares and rods [290, 291], dimers and ellipses [292], and others [293, 294]. It would be interesting to build network representations of these systems, examine how different grain geometries affect network organization, and investigate how that organization relates to the mechanical properties of a system [295]. It seems particularly important to develop an understanding of which (quantitative and qualitative) aspects of network structure depend on features of grains (such as shape, polydispersity, friction, cohesiveness, and so on [27, 164, 168, 169, 176, 180, 210]) and which are more universal.

One can also consider defining particulate networks in a variety of other ways. For example, when determining edges and edge weights, one can examine the tangential (rather than, or in addition to, the usual normal) component of the force between two grains. Such extensions may facilitate increasingly detailed investigations into a packing’s organization [166]. It may also be useful to retain information about both the magnitude and direction of forces when defining edges. One may even wish to construct signed networks, for which edges can take either positive or negative values, thereby conveying further information about the relationship between nodes. In such studies, one can perhaps take advantage of advancements in community-detection techniques, such as by using signed null models [296, 297]. Additionally, as we discussed in Section 3.3, particle-property networks [114, 116, 269] and networks constructed from particle-displacement information [28, 115, 117, 162] are other informative ways to build networks for particulate systems. One can also construct edges (and determine edge weights) by incorporating information about inter-grain relationships based on similarities in particle properties such as orientation [298] (see Fig. 23), coefficient of friction [299], or size [300]. Constructing networks whose edges are determined or weighted by inter-particle similarities may be particularly useful for achieving a better understanding of mesoscale physics in polydisperse packings, which are thought to depend on the spatial distributions of particles of different types [301]. A perhaps non-intuitive choice is to use a bipartite representation of a granular network, such as the approach used in [302].

The above choices for network construction give a grain-centric view of the physics of particulate materials. One can also consider edge-to-node ‘duals’ of such networks to provide a contact-centric perspective. In a contact-centric approach, one treats contacts between physical objects as nodes and the objects themselves as edges between contacts. (Compare this idea to the notion of a line graph

10 Other ideas that are worth considering include memory networks [285], adaptive networks [49], and various representations of temporal networks [286].
Fig. 23. Constructing a network representation of collagen-fibre alignment. We show quantitative polarized light images (QPLIs) of the human cervical facet capsular ligament taken both before and during loading. These data were collected as part of a study of collagen-fibre realignment following a mechanical stretching of the material. Such realignment is a mechanical process that commonly leads to both acute and chronic neck pain, particularly following car accidents. QPLIs were used to generate pixel-wise collagen alignment maps in the selected regions of interest (ROI) of anomalous realignment (AR; red box) and normal realignment (NR; blue box) at rest (time point $t = 1$), at the onset of AR (time point $t = T$), and in between these two points in 1-second increments. Rectangular ROIs were defined using the upper-left and lower-right fiduciary markers as the common information between different time points. Each $3 \times 3$ pixel window in an ROI is a node in a network. We show representative $3 \times 3$ pixel windows in the resting and stretched states, along with a corresponding demonstration of alignments. For each node, we calculate the alignment angle, and we use a measure of similarity between alignment angles to weight the edges in the network. In this way, weighted adjacency matrices, with nodes numbered spatially, illustrate a pairwise similarity in alignment angles between ROIs. We reproduced this figure, with permission, from [298].

[303] of a network $G$.) A contact-centric network approach was used recently in the study of nanorod dispersions [304, 305]. Contacts between rods were treated as nodes, and the effective conductance of the rod was treated as a weighted edge. The treatment of a grain or other physical object as an edge rather than as a node is also a particularly appealing way to describe networks of fibres in both human-made and natural systems. Recent examples of such fibre networks that have benefited from network analysis include collagen networks [298, 306] (see Fig. 23), fibrin networks [307], and axonal-fibre networks [57, 308].

Another way to study granular systems (especially porous ones) is to consider a network constructed from the pore space [309], in which pores (i.e., empty volumes between contacting grains) are nodes and throats (i.e., flow pathways that connect pores) are edges (which can be weighted in various ways). Conveniently, there are several methods to precisely determine pores and grains [310]. Studying pore networks is a common way to examine flow through porous materials or to understand the responses of granular materials to external stresses, but only more recently have such networks been studied explicitly from a network-science perspective. See [311–315] for some recent examples, and see [316] for a study of force chains in porous media.
4.1.2 Multilayer networks. When considering different ways to construct a network, it is also natural to ask whether it is beneficial to combine two or more approaches in an integrated way to more accurately (and elaborately) represent a particulate system as a network. Ideally, doing network analysis using a more complicated representation will also lead to improved physical understanding. One way to incorporate heterogeneous node types, multiple types of relationships between nodes, and time-dependence into network representations is with multilayer networks [36, 135, 137]. Recall that [113] (see Section 2.2.7) studied one type of multilayer community structure in a compressed granular material. Another type of multilayer network that may be useful for the study of particulate systems is a multiplex network, in which one can encode different types of relationships between nodes in different layers of a network. For example, one can construct a multiplex network in which two particles are linked both by normal force-weighted contacts and according to another relationship (such as the tangential force between them, their contact angle, or by a measure of similarity of one or several particle-properties, as discussed in Section 3.3.3). One can also envision using multilayer networks to study particulate systems with multiple types of particles. For example, if a system consists of particles of different shapes or sizes, one possibility is that each particle of a given shape (or size) is in a different layer, intralayer edges represent interactions between particles of the same type, and interlayer edges represent interactions between particles of different types. Another possibility is to let each layer represent a time window, perhaps with intralayer edges representing a mean interaction strength during that window. The study of multilayer networks is one of the most popular areas of network science, and we expect that such networks will be very illuminating for studies of particulate matter.

4.1.3 Annotated graphs. The network-construction techniques that we have been discussing throughout this review represent data in the form of an adjacency matrix (for an ordinary graph) or an adjacency tensor (for a multilayer network) [36, 136, 137]. However, it may be desirable to encode not only relationships between grains, but also features of the grains and/or their interactions. One option is to use annotated graphs (one can also use multilayer networks) to encode inter-node relationships, individual-node properties, and individual-edge properties [317, 318]. One can use annotated graphs—also sometimes called labelled graphs or tagged graphs [319]—to study properties of interaction patterns (such as force-weighted contacts) that may be driven by local particle properties (e.g., size, shape, spatial position in a material, membership in cycles or force chains, and so on). Available tools for annotated graphs include clustering techniques in the form of SBMs that combine information from both connectivity and annotations to infer densely-connected sets of nodes [317, 318].

4.1.4 Beyond pairwise interactions. It is also desirable to explicitly examine interactions between three or more particles, rather than restricting one’s study to representations of pairwise interactions. We discussed this idea briefly in Section 2.2.10 in the context of simplicial complexes [320], and we note that one can also encode edges among three or more nodes by using hypergraphs [29].

4.1.5 Physically-informed network calculations. In addition to exploring different choices of how to construct particulate networks, it is also important to consider ways to generalize the tools of network science to incorporate physical constraints [27, 37]. A natural way to begin is to build null models of spatially-embedded graphs that obey simple mechanical constraints like force balance [321]. One can also develop network diagnostics and methods that incorporate physical geometry, spatial embedding, and latent spatial structure of networks [37]. For such investigations, we expect that ideas from the
study of random geometric graphs, random neighbourhood graphs, and their extensions will be helpful [21, 351, 352]. It will also be useful to employ kinetic approaches, such as the one in [322], and to incorporate ideas of flow and other forms of dynamics into community detection [122, 323]. Additionally, it is desirable to develop methods to more explicitly characterize the relationship between a network’s structure and the geometry of a space in which it is embedded [37, 324, 325] (as well as latent geometry) and to use such techniques to better understand the plurality of packings that are consistent with a single force distribution via a force-network ensemble, which is the space of allowed force configurations for a fixed contact geometry [202–205, 326]. We also point out that a spatial embedding can induce correlations between network diagnostics [21], and it is therefore important to develop a better understanding of the extent to which this occurs in networks that arise from particulate systems.

4.2 Beyond granular materials

Although we have focused primarily on network analysis of the canonical type of granular systems, which consist of discrete, macroscopic particles that interact via contact forces, one can potentially also use network-based approaches to characterize granular materials with more complex interactions as well as soft materials more broadly [327]. As reviewed recently in [6], these materials include colloids, glasses, polymers, and gels. In these systems (and others) the particles (or other entities) can interact via various (and often complicated) means. For example, system components can have attractive and/or repulsive long-range interactions [329], can be cohesive [330], and/or can interact with one another via chemical gradients [331, 332], electric charges [333–335], or through other molecular or mechanical processes [336]. In each of these cases, interaction strengths can yield edge weights, either as the sole relationship studied between entities in a graph or as one of several inter-entity relationships in a multilayer network.

One particularly interesting avenue for future work may be to study polymer and fibre networks [337, 338], which are important in biological systems. For example, in biopolymer assemblies, cross-linking can glue filaments together into large-scale web-like structures (e.g., as in the cytoskeleton of a cell) [339]. Such cross-linked actin filaments are critical for cellular function and integrity [340], and it is thus important to understand the structural organization of these kinds of networks, their mechanical properties, and how force transmission is regulated in them [341–343]. Indeed, there has already been some work examining network representations of gels and polymers, and employing graph-theoretic analyses to quantify the structural properties of these systems (see, e.g., [344–349]). One can also use network analysis to study systems at smaller spatial scales. Because traditional network-based approaches are agnostic to the physical scale of a system, off-the-shelf calculations and algorithms are directly portable to microscale, nanoscale, and even smaller-scale systems [327, 350]. However, despite the technical portability of common tools, the investigation of much smaller-scale systems should benefit from the extension of classical network-based tools in ways that incorporate additional underlying physics. For example, we have described extensions of network tools to assimilate ideas and constraints from classical physics (e.g., spatial embeddedness) [27, 129]. One can also consider employing ideas that take into account principles from quantum physics [353, 354] or other areas. The study of quantum networks is a new and exciting area of network science, and there are many opportunities for progress [355].

---

11 One can also examine particulate networks of hard materials that admit Hamiltonian descriptions [328].
4.3 Implications for material design

As is the case with mathematics more generally, network analysis gives a flexible approach for studying many systems, because it is agnostic to many details of their physical (or other) nature [29, 86, 356]. Such versatility supports the application of network-science techniques to both living and non-living materials to examine the architectures and dynamics of these systems and to gain insights into relationships between structure and function. The tools and approaches of network science also have the potential to inform the design of new materials. For example, it should be possible to use network theory (e.g., via the tuning of a system’s network architecture) to provide guidance for how to engineer a material to exhibit specific mechanical, electrical, or other properties. Material design has become increasingly popular with recent advances in the study and development of metamaterials [357–359]. Metamaterials can take advantage of precisely-defined component shapes, geometries, orientations, arrangements, and connectivity patterns (rather than specific material and physical characteristics of individual units) to produce tailored mechanical [360], acoustic [281, 359, 361–363], and electromagnetic [357, 364, 365] properties. The control of a single unit or component is relatively straightforward, but the question of how to link many components in ways that yield complex material properties is a very challenging one. Approaches that use ideas from network science have the potential to offer guidance for construction patterns of material units that support desired bulk properties.

There are likely many ways to use network-based approaches to inform the design of new materials. One reasonable possibility is to employ evolutionary and genetic computer algorithms [366, 367] and other tools from algorithmic game theory [368]. For example, the combination of multi-objective functions and Pareto optimality [369] can offer a targeted path through the space of possible network architectures (i.e., through a network morphospace [370]). If exact simulations are not computationally tractable, one can use machine-learning techniques to offer fast estimates of material properties and behaviour [371]. One can perhaps begin with a single material network structure that is physically realizable and then rewire the initial network architecture with a cost function that drives the system towards an arrangement that enhances a desired property. One can perform such a rewiring explicitly along Pareto-optimal fronts using a set of rewiring rules that preserve physical constraints and laws. This approach, which builds on prior applications to other types of spatially-embedded networks [372–374], selects for physically-feasible network designs that purposely maximize specific properties. A network analysis of these ‘evolved’ (and evolving) networks may help elucidate relationships between structural features of system architecture (e.g., values of a specific network diagnostic) and material properties (e.g., stability under load), providing a link between structure and function. Such an evolutionary-design approach can complement recent efforts to identify optimal shapes with which to construct a packing [375, 376], to design rules with which to pack those shapes [377, 378], and to construct ‘allosteric materials’ with specific functionalities via evolution according to fitness functions [379].

One set of problems for which network-based tools may be useful are those related to rigidity and mechanical responses of disordered material networks, which have been studied previously using a rigidity-percolation framework [170, 172, 177, 380]. In terms of material design, a particularly interesting line of future work may be to use network analysis in conjunction with methods such as tuning-by-pruning, in which individual bonds are selectively removed from a disordered material network to engineer a specific property into a system [381]. For example, beginning with simulated, disordered spring networks (derived from jammed particle packings), Goodrich et al. [381] used tuning-by-pruning to design networks with different ratios of the shear modulus to the bulk modulus. Motivated by allosteric responses in proteins, [382] developed an approach that allows careful control of local mechanical responses in
disordered elastic networks. In particular, with the removal of very few bonds, one can tune the local strain in one region of a network via a response to an applied strain in a distant region of a system. Driscoll et al. [383] studied three different spring networks and continuously tuned their mechanical rigidity (measured by different parameters, such as the distance above isostaticity) to examine the effect of such tuning on material failure. They observed that for a fixed amount of disorder (which the authors measured using a scalar, following the approach in [384]), the width of the failure zone in a stressed material increases as the rigidity decreases. Recently, [385] studied a model of amorphous networks that incorporates angle-bending forces, and employed pruning-based methods to design auxetic metamaterials. In light of these findings, it is natural to ask what network quantities and methods may be related to the above types of global or local mechanical properties, and whether they can inform which bonds to remove (or rearrange [224]) to invoke particular types of functional responses.

In developing a network-based approach for designing and building new materials, it is desirable to capitalize on the ability of network analysis to quantify multiscale structures (with respect to space, time, and network architectures) in a wide variety of systems (regardless of the exact details of their composition). For example, network analysis has revealed mesoscale architectures that are often crucial for determining material properties in disordered media, but such heterogeneities also appear to be important for biological materials, including networks of collagen fibres [298, 386], tissues and tendons [387, 388], muscle fibres [389], and axonal fibres [56, 390]. Tools from network science should be useful (and, in principle, flexible) for designing non-trivial structural organizations that yield desired material functions. One can imagine using a network-theoretic framework to design localized, mesoscale, and/or system-level properties, to design and manipulate human-made and natural (including biological) materials, and to precisely control both static and dynamic material properties.

5. Conclusions

Network science is an interdisciplinary subject—drawing on methods from physics, mathematics, statistics, computer science, social science, and many other disciplines—that has been used successfully to help understand the structure and function of many complex systems. Much recent work on networks has yielded fascinating insights into granular materials, which consist of collections of discrete, macroscopic particles whose contact interactions give rise to many interesting behaviours and intricate organization on multiple spatial and temporal scales. These insights have increased scientific understanding of the structure and dynamics of the heterogeneous material architecture that manifests in granular matter (as well as network-based approaches to quantify such architecture) and the response of granular systems to external perturbations such as compression, shear, tapping, and tilting. In this article, we have reviewed the increasingly fertile intersection of network science and granular materials. Future efforts should help provide a better understanding of the physics of particulate matter more generally, elucidate the roles of mesoscale interaction patterns in mechanical failure, inform the design of new materials with desired properties, and further scientific understanding of numerous important problems and applications in granular physics, soft-matter physics more generally, and even biophysics.

Acknowledgements

We thank Alejandro J. Martínez, Ann E. Sizemore, Konstantin Mischaikow, Jen Schwarz and an anonymous referee for helpful comments. We also thank our collaborators, students and mentors, who have shaped our views on the subjects of this review.
Funding

KED is grateful for support from the National Science Foundation (DMR-0644743, DMR-1206808) and the James S. McDonnell Foundation. LP is grateful to the National Science Foundation for a Graduate Research Fellowship. DSB is grateful to the Alfred P. Sloan Foundation; the John D. and Catherine T. MacArthur Foundation; the Paul G. Allen Foundation; and the National Science Foundation (PHY-1554488). The content is solely the responsibility of the authors and does not necessarily represent the official views of any of the funding agencies.

REFERENCES


Upper Saddle River, NJ, USA: Prentice-Hall, Inc.


