Non–universality of elastic exponents in random bond–bending networks

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We numerically investigate the rigidity percolation transition in two–dimensional flexible, random rod networks with freely rotating cross-links. Near the transition, networks are dominated by bending modes and the elastic moduli vanish with an exponent $f = 3.0 \pm 0.2$, in contrast with central force percolation which shares the same geometric exponents. This indicates that universality for geometric quantities does not imply universality for elastic ones. The implications of this result for actin–fiber networks is discussed.

In contrast with ordered crystals, disordered materials typically exhibit a complex connection between their collective mechanical properties and the underlying interactions between their constituent elements \cite{2}. In this Rapid Communication, we study this connection between effective elastic theories of a flexible rod network in two dimensions and the fundamental elastic properties of individual rods, along with statistical measures of the network connectivity. We focus on cross-link densities to explore the break-down of collective rigidity of the network. This vanishing of the static shear modulus in related systems \cite{1, 3, 4, 5, 6} has been termed the rigidity percolation transition in analogy to the better understood scalar, or conductivity, percolation transition \cite{6}. Previously \cite{4}, we investigated the internal deformation field of a 2d flexible rod network away from the critical point of rigidity percolation, and found a cross-over from affine to nonaffine deformation upon changing the mean density between cross links in the network and/or the inherent flexibility of individual rods, which was also independently corroborated by another group \cite{5}.

The rigidity percolation transition has been previously studied numerically in a number of model, disordered systems with variations in both the nature of the links (bonds) and the disorder of the network. While the rigidity transition is first order on the Bethe lattice \cite{11} and in random networks of infinitely rigid rods \cite{12}, in all other random network models it is continuous and characterized by a diverging length scale over which the material acquires a finite, static shear modulus. From the outset, it is important to distinguish between two different classes of physical observables when discussing the long length scale physics at the transition: (i) geometric quantities which describe the fractal structure of the percolating rigid cluster at the transition; and (ii) elastic properties of the material near the percolation point. In the first category are the exponents $\nu$ and $\beta$, which describe the divergence of the correlation length as one approaches the transition and the probability that a link will be part of the percolating cluster, respectively. In the second category is $f$, which describes the power law dependence of the system’s elastic moduli upon the approach to the percolation transition from the rigid phase (see Eq. 3).

In lattice models, the control parameter, as for scalar percolation, is the probability of the presence of a link in the diluted lattice. It appears that the nature of disorder is a relevant variable, since for purely central forces between network nodes, site and bond disorder exhibit quantitatively different scaling regimes near the critical point \cite{1, 3}. In addition, a third universality class has been postulated in the so-called “bond–bending” model, where bending lattice edges and rotation at vertices cost energy \cite{1, 3, 4, 5}. This suggests that the introduction of a bending modulus in the model is a relevant perturbation at the transition.

Pertinent to the current work are the simulations of Latva-Kokko et al. \cite{1, 2}. These investigations, departing from previous work, turned to off–lattice simulations. They constructed two–dimensional random rod networks and applied a topological approach to investigate the rigidity percolation point in random rod networks with Hookean central forces and a bending modulus. The introduction of a bending modulus is vital at this stage since the random spring network that is not prestressed has zero–frequency deformation modes at any finite density of links, and is thus always nonrigid \cite{14}. Latva-Kokko et al. considered two variants of their model of flexible rods distinguished by the constraint forces imposed at a cross link: one with cross links that fix the angle between intersecting rods by applying local constraint torques, and freely rotating bonds at cross links in the other. In both cases they found that the geometric exponents are consistent with those of the rigidity percolation transition in a diluted lattice with central forces. This result suggests that the introduction of bending forces are not relevant perturbations at the critical point in the following restricted sense: the exponents associated with length scale and geometry of the spanning, rigid cluster appear to be universal. We use this result later. Their approach, however did not allow them to investigate the scaling exponent for the elastic quantities of their model.

This Rapid Communication presents a study of the elastic properties of the rod network identical to the model system studied by Latva-Kokko et al. We, however, concentrate on the mechanical aspects of the network near the transition. It is important to note that this work relates to models of semi–flexible polymer networks with freely rotating bonds \cite{14}. This detail of the...
model distinguishes it from its other close antecedent, the bond-bending model of S. Feng et al. That lattice–based model has been particularly well described by both numerical simulation and real-space renormalization group techniques \[16\]. Our model differs in that the two filaments that cross at each node contribute independently to that node’s bending energy; there is no energy cost for relative rotation between rods. Previous lattice calculations suggest that the scaling of the elastic constants near the transition depend on such details of the network; with this in mind, we seek herein to explore the rigidity collapse of sparse actin networks.

The principal result of this communication is that the mechanical properties of the flexible rod network at the rigidity percolation transition are distinct from previously investigated models of either lattice–based bond-bending networks or central force networks. This is true despite the fact that the scaling of the size of the percolating cluster and its fractal geometry (as determined by Latva-Kokko et al. \[6\]) are identical to that of central force networks. This point highlights the physical independence of geometric/topological exponents and the elastic exponents of the network. We propose that whilst some degree of universality in the elastic properties of disordered systems may exist, such universality classes are smaller and more numerous than for geometric properties; that is, models with the same geometric exponents \(\nu\) and \(\beta\) may have distinct elastic exponents \(f\). Our model system below provides a concrete example of this claim. We suspect that the underlying cause of this non–universality is that while the rigidity transition itself depends on large scale topological properties of the network that are insensitive to the local details of stress transmission, the moduli of the rigid network near the transition are sensitively determined by only a vanishingly small set of paths for stress propagation. The moduli thus depend on local details of how stress is transmitted through the small set network nodes along the weak points of this path. Different force laws at e.g. cross links can therefore have profound effects on the way in which the moduli vanish at the transition.

We numerically evaluated the elastic moduli of random networks of rods at \(T = 0\), with an energy \(\delta H\) per unit length \(\delta s\) given by

\[
\frac{\delta H}{\delta s} = \frac{\mu}{2} \left( \frac{\delta l}{\delta s} \right)^2 + \frac{\kappa}{2} \left( \frac{\delta \theta}{\delta s} \right)^2
\]

The first term on the right hand side describes an elastic restoring force for changes in relative length of the rods, \(\delta l/\delta s\), with a spring modulus \(\mu\). Bending by an angle \(\delta \theta\) incurs an energy cost given by the second term, where the bending modulus \(\kappa\) is the same as in the wormlike chain model \[13\]. Cross linked rods are coupled by imposing the same coordinates at intersections.

The simulation method is described in detail elsewhere \[17\] (where we also discuss behavior away from the transition), but in brief: rods of length \(L\) are deposited with random orientation and position into a two-dimensional shear cell of dimensions \(W \times W\). Each intersection is identified as a cross link, the mean distance between which (as measured along a rod) is denoted \(l_c\), so that the mean number of crosslinks per rod is \(L/l_c - 1\). Deposition continues until the required cross linking density \(L/l_c\) has been reached. The system Hamiltonian is constructed from \[1\], and the mechanical equilibrium configuration found by the preconditioned conjugate gradient method under the constraint of an applied shear or uniaxial strain. An example is given in Fig. \[1\].

![Fig. 1](image-url) (Color online) The energy density for a network at the transition \(L/l_c \approx 5.933\) under an applied shear strain. The line thickness is proportional to the logarithm of the energy density per unit length. Apparent stressed ‘dangling’ ends are artefacts of numerical noise and make no significant difference to the measured quantities discussed below. The calibration bar shows what proportion of the energy is due to stretching. For clarity, a small \(W = 7.5L\) shear cell is shown.

There are four lengths in the problem: the system size and rod length \(W, L\) respectively; the mean distance between cross links \(l_c\) and a length characterizing the flexibility of the rods, \(l_h = \sqrt{\kappa/\mu}\). We take \(L/l_c\) as our dimensionless measure of density rather than \(q = NL^2\) with \(N\) the number of rods per unit area, as in \[3, 7, 14\]. It is straightforward to convert between the two measures using the expression derived in the Appendix. Rigidity percolation occurs at \((L/l_c)_\text{trans} \approx 5.933\). For \(L/l_c\) slightly above this critical point, where \(\varepsilon = (L/l_c)/(L/l_c)_\text{trans} - 1 \sim 0^+\), we write the shear modulus \(G\) as

\[
G = \frac{\kappa}{L^2} f \left( \varepsilon, \frac{l_c}{L}, \frac{W}{L} \right)
\]

with a similar expression for the Young’s Modulus, \(Y\). For \(\varepsilon > 0\), and sufficiently large system sizes \(W\), both \(G\) and \(Y\) depend only on \(\varepsilon\) and \(l_c/L\). Fig. \[2\] shows \(G\) and \(Y\).
TABLE I: Quoted \( f \) for 2d uncorrelated networks

| Model                        | \( f \)  | Ref. |
|------------------------------|---------|------|
| Site percolation             | 1.35 ± 0.06 | [4, 5] |
| Central force percolation    | 3.57 ± 0.3  | [4, 5] |
| Bond–bending model           | 3.96 ± 0.04 | [4, 8, 20] |
| This work                    | 3.0 ± 0.2  | n/a  |

versus \( \varepsilon \) for systems where this large–\( W \) limit has been reached. Both moduli vanish continuously as \( \varepsilon \to 0^+ \), confirming both the nature and location of the transition as claimed in [6, 19], to within our data resolution [18].

Fig. 2 shows that \( GL^3/\kappa \) is independent of \( l_b/L \) for sufficiently small \( \varepsilon \), i.e., \( G \) is independent of \( \mu \) near the transition. This suggests that the transition is dominated by bending modes, and we can infer that \( G \) would vanish if there were only central force terms in agreement with [14]. We can confirm the dominance of bending modes at the transition by measuring the proportion of stretching energy for increasing \( W \). As shown in inset of this figure, this fraction is always small, \(<4\%\), and may vanish as \( W \to \infty \), as suggested by the figure and in agreement with both [14] and our observation that \( G \propto \kappa \).

By fitting the data to the functional form

\[
G = A\varepsilon^f(1 + B\varepsilon) \tag{3}
\]

we find \( f = 3.0 \pm 0.2 \), consistent with [2], where the error estimate gives the spread of values when the fitting procedure is repeated over different data subsets. \( B \) is always \( O(1) \), suggesting [3] is a ‘sensible’ choice. This is compared to known values in Table I. The only possible equality is with central force bond percolation; however, since the modulus near the transition is dominated by bending forces, we do not believe that this near agreement is meaningful. The fit for \( G \) can be made to agree with the Young’s modulus data by simply rescaling \( A \to A’ \) with \( f \) and \( B \) fixed, as shown in Fig. 2 for \( l_b/L = 0.006 \). \( A’/A = 2.7 \pm 0.2 \), so that the Poisson ratio \( \sigma = Y/2G - 1 = 0.35 \pm 0.1 \) at the transition, consistent with the claimed ‘universal’ value of \( \frac{1}{3} \) [14].

Independent confirmation of \( f \) can be found by placing the system at the critical point \( \varepsilon = 0 \) and considering the variation of \( G \) with \( W \) [4, 8, 9]. Scaling arguments suggest that \( G \sim W^{-f/\nu} \) for sufficiently large \( W \), where \( \nu \) is the exponent describing the divergence of the correlation length, which is known for these networks to be in the range of \( \nu = 1.17 \pm 0.02 \) [6]. We attempted to fit our data to the same functional form used in [6], but the precision with which we can extract the exponent using this approach is not sufficient to distinguish our results from either central force or bond–bending models. The number of networks for each attempted \( W \) ranged from 400 to 2000, comparable to lattice models which give much smaller errors. Our lack of precision is likely due to the random nature of our networks, in which the distance between crosslinks varies continuously down to zero so that the strength of coupling between connected nodes can vary greatly, introducing a significant additional noise source into the problem.

In summary, we have shown that the elastic moduli of flexible random networks with freely rotating crosslinks scale as \( \varepsilon^f \) near to rigidity percolation. The modulus itself is dominated by bending modes near the transition and vanishes with an exponent \( f = 3.0 \pm 0.2 \). This exponent differs from previous reported exponents for models that include bond bending. Our introduction of bending forces, motivated as it was by modelling cross linked F-actin networks, differs in detail from previous work.
Our results suggest that, in contrast to the geometric exponents previous reported for this system, the modulus exponent is more highly model dependent. To address the experimental implications of these results for physical actin networks, we note that the continuous transition with true scale invariant behavior exists at only zero temperature. However, zero temperature phase transitions can have experimental consequences at finite temperature [22] as long as it is possible to experimentally access the critical region. We expect that in sparse actin networks, we note that the continuous transition with true scale invariant behavior exists at only zero temperature. We expect that in sparse actin networks, we note that the continuous transition with true scale invariant behavior exists at only zero temperature. We expect that in sparse actin networks, we note that the continuous transition with true scale invariant behavior exists at only zero temperature. 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