Interpretation of the Vibrational Spectra of Glassy Polymers Using Coarse-Grained Simulations

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ABSTRACT: The structure and vibrational density of states (VDOS) of polymer glasses are investigated using numerical simulations based on the classical Kremer–Grest bead–spring model. We focus on the roles of chain length and bending stiffness, the latter being set by imposing three-body angular potentials along chain backbones. Upon increasing the chain length and bending stiffness, structural reorganization leads to volumetric expansion of the material and buildup of internal stresses. The VDOS has two dominant bands: a low-frequency one corresponding to inter- and intrachain nonbonding interactions and a high-frequency one corresponding principally to vibrations of bonded beads that constitute skeletal chain backbones. Upon increasing the steepness of the angular potential, vibrational modes associated with chain bending gradually move from the low-frequency to the high-frequency band. This redistribution of modes is reflected in a reduction of the so-called Boson peak upon increasing chain stiffness. Remarkably, the finer structure and the peaks of the high-frequency band, and their variations with stiffness, can, for short chains, be explained using an analytical solution derived for a model triatomic molecule. For longer chains, the qualitative evolution of the VDOS with chain stiffness is similar, although the distinct peaks observed for short chains become increasingly smoothed out. Our findings can be used to guide a systematic approach to interpretation of Brillouin and Raman scattering spectra of glassy polymers in future work, with applications in polymer processing diagnostics.

I. INTRODUCTION

Raman spectroscopy can detect vibrational and electronic properties of materials over a broad range of temperatures and pressures and is a well-established and widely used non-destructive measurement technique. Comprehensive predictive models for Raman and Brillouin spectra are important for many applications involving amorphous carbon-based materials, from nanotechnology to polymer reaction engineering. Of particular interest is emulsion polymerization, a common manufacturing route for many rubbers and plastics. The complexity of this process hinders characterization of product quality by traditional methods, and it is increasingly being probed by Raman spectroscopy.

The vibrational density of states (VDOS) of solids is the main input for the prediction of the Raman and Brillouin scattering spectra. For glasses, the Shuker–Gammon formula gives the Raman intensity as a function of the VDOS as

\[ I(\omega) = \frac{n(\omega) + 1}{\omega} C(\omega) D(\omega) \]  

(1)

where \( n(\omega) + 1 \) is the Bose–Einstein occupation factor, \( D(\omega) \) is the VDOS, and \( C(\omega) \) is the photon–phonon coupling coefficient. Since \( C(\omega) \) is a simple function of frequency, possibly quadratic, \( I(\omega) \) is proportional to \( D(\omega) \). It is clear that most of the structure of the Raman spectrum is directly related to the \( D(\omega) \) spectrum. While the VDOS of crystals can be obtained by a straightforward exercise in Fourier analysis, the same problem for amorphous solids, such as glasses, is analytically intractable and presents a rich phenomenology. This phenomenology is yet more complex when the building blocks are polymer chains, which, in the disordered glassy state, can have a considerably larger variety of conformations.

There have been numerous studies into the vibrational properties of polymeric systems starting from theoretical determinations of the single-chain backbone vibrational spectra in seminal works by Kirkwood and Pitzer, followed by the powerful combination of Wilson’s GF method with group theory by Higgs. These methods are not applicable to polymer glasses, however, where the chain conformation does not possess any periodicity that can allow the application of group theoretical methods. Further advances in numerical techniques have focused on reducing the computational time of the diagonalization problem.

While signatures of individual monomers and their constituent bonds are very well characterized in the vibrational bands of highest energy in the spectrum, the relation between coarse-grained polymer structures and vibrational properties in the low-frequency part of the spectrum is relatively unexplored. In the contemporary literature, the use of coarse-grained systems as model materials for studying the vibrational properties of amorphous solids has become a standard...
In this direction, coarse-grained simulations based on the classical Kremer–Grest model\(^\text{12}\) for bead–spring polymers can enable direct calculation of the VDOS. With a suitable procedure for coupling the VDOS to the Raman spectra,\(^\text{18}\) such numerical approaches will be able to offer a systematic approach to linking vibrational properties to coarse-grained structures for polymers of arbitrary length and monomer–monomer interactions.

Here, we report the structural properties and the VDOS for coarse-grained polymer glasses as functions of the chain length and the chain bending stiffness. We identify clear trends in the vibrational spectra that derive from microstructural rearrangements as the chain length and chain stiffness increase. Through these quantities, it will be possible in future work to make predictions about how the experimentally observed vibrational spectra will evolve during the course of an emulsion polymerization, for example, guiding the development of noninvasive industrial process monitoring techniques. This work can further serve as the basis for quantitative understanding and modeling of Raman and Brillouin spectra at the atomistic level, particularly by coupling to atomistic simulation techniques.

In the following, we first describe the numerical method used and then go on to study the structural and volumetric changes as functions of varying chain length and stiffness. We then analyze the VDOS as a function of chain length and stiffness, providing a mechanistic interpretation informed by an analytical argument.

**II. SIMULATION DETAILS**

Our model uses a coarse-graining approach that treats polymer chains as linear series of monomer “beads” on an elastic string. In a harmonic approximation, monomeric scale physics dominates the region of the VDOS of interest to this work and indeed governs the viscoelastic response of the material.\(^\text{22}\)

For each bead in the system we use LAMMPS\(^\text{23}\) to solve the Langevin equation

\[
\frac{m}{d t} \frac{d r}{d t} = -\frac{m}{\zeta} v - \frac{d U}{d r} + f_b(t)
\]

for uniform beads of mass \(m\) and velocity \(v\), coefficient of friction \(m/\zeta\), and random forces \(f_b(t)\) satisfying \(f_b(t)f_b(t') = 2mk_B T \delta(t-t')/\zeta\). Beads interact with each other through a potential \(U\), given by the Kremer–Grest model\(^\text{12}\) with the addition of angular potentials that impose bending constraints on triplets of three consecutive beads along the chain backbones. Overall, the model for the potential energy \(U\) comprises three terms: [i] A truncated and shifted Lennard–Jones (LJ) potential of form

\[
U_{\text{LJ}}(r) = 4\varepsilon_{\text{LJ}} \left[\left(\frac{\sigma}{r}\right)^{12} - 2\left(\frac{\sigma}{r}\right)^{6}\right]
\]

acting between all bead pairs within a cutoff range \(r_c = 2.5\sigma\), where \(r\) is the bead–bead separation, \(\sigma\) is the zero-crossing distance for the potential, and the prefactor \(\varepsilon_{\text{LJ}}\) sets the LJ energy scale. Setting \(\frac{dU_{\text{LJ}}(r)}{dr} = 0\) leads to an energy minimum and corresponding LJ rest position at \(2^{1/6}\sigma\). The LJ potential effectively acts as an excluded volume, as illustrated in Figure 1. [ii] A finitely extensible nonlinear elastic (FENE) potential acting between sequential bead pairs along each linear chain

\[
U_{\text{FENE}}(r) = -0.5\varepsilon_{\text{FENE}} R_0^2 \ln \left[1 - \left(\frac{r}{R_0}\right)^2\right]
\]

where \(R_0\) is the maximum FENE bond length and \(\varepsilon_{\text{FENE}}\) is the bonding energy scale. Adjacent beads along polymer chains have an overall interaction that represents the sum of the Lennard-Jones and FENE potentials, giving a rest position for bonded beads (obtained by setting \(\frac{d}{dr}(U_{\text{LJ}}(r) + U_{\text{FENE}}(r)) = 0\)) as \(\approx 0.96\sigma\) for the parameters used throughout this work. This discrepancy relative to the LJ rest length gives sufficient bidispersity to suppress crystallization throughout, and we set \(\varepsilon_{\text{FENE}}/\varepsilon_{\text{LJ}} = 30\). (iii) An energy associated with chain bending, given by

\[
U_{\text{bend}}(\theta) = \varepsilon_{\text{bend}} [1 - \cos(\theta - \theta_0)]
\]

for energy scale \(\varepsilon_{\text{bend}}\). The angle \(\theta\) is formed between three consecutive beads (a triplet) along the length of the linear chains (Figure 1), and the characteristic rest angle is \(\theta_0 = 109.5^\circ\). The resistance to bending of the polymer chains is thus set by \(\varepsilon_{\text{bend}}\), which is related to the persistence length \(l_p\) of the chain via the standard relation \(l_p = \varepsilon_{\text{bend}}/\kappa_b T\).

The relative importance of the three potentials in setting the overall structure and dynamics of the polymers is determined by their prefactors \(\varepsilon_{\text{LJ}}, \varepsilon_{\text{FENE}}, \text{ and } \varepsilon_{\text{bend}}\). Since each of the potentials has a different form, it is difficult to compare the values of these prefactors directly. In order to render the different interaction strengths more comparable, therefore, we find it convenient to take a harmonic approximation about the rest position of each potential and consider the resulting spring constants \(k\). We find \(k_{\text{LJ}} \approx 57.1\varepsilon_{\text{LJ}}/\sigma^2, k_{\text{FENE}} \approx 32.7\varepsilon_{\text{FENE}}/\sigma^2,\) and \(k_{\text{bend}} = \varepsilon_{\text{bend}}/\sigma^2\). To characterize our systems, we use two control parameters \(k_{\text{bend}}/k_{\text{LJ}}\) and \(k_{\text{bend}}/k_{\text{FENE}}\) that compare the bending stiffness to the LJ and FENE bond strength, respectively. The strength of FENE bonds is fixed such that \(k_{\text{FENE}}/k_{\text{LJ}} \approx 17.2\) throughout (recalling that \(\varepsilon_{\text{FENE}}/\varepsilon_{\text{LJ}} = 30\)). We explore bending stiffnesses in the range \(k_{\text{bend}}/k_{\text{LJ}} = 0 \rightarrow 20\). A sketch of two interacting polymer chains is shown in Figure 1, highlighting the angle \(\theta\) on which \(U_{\text{bend}}\) acts as well as the rest positions for LJ (\(2^{1/6}\sigma\)) and FENE (\(0.96\sigma\)) interactions.

With reference to fundamental units of mass \(\mu\), length \(d\), and energy \(\varepsilon\), we set \(\sigma = 1, R_0 = 1.5, m = 1, \) and \(\varepsilon_{\text{LJ}} = 1\), giving a time unit of \(\tau = \sqrt{\mu\sigma^2/\varepsilon_{\text{LJ}}}\), and we set \(\xi = 100\tau\). The system volume \(V\) has units \(d^3\). A dissipative time scale emerges as

![Figure 1. Sketch of simulated polymer system showing two interacting chains (blue and pink). Shown are the LJ rest position of \(2^{1/6}\sigma\) between nonbonded beads, the FENE rest position of \(0.96\sigma\) between beads along a single linear chain, and the angle \(\theta\) formed between three consecutive beads (forming a triplet) along chains.](Image)
and a thermal time scale emerges as $m\sigma^2/\xi_{FL}$, and the state of our system, i.e., whether it is in the melt or glassy state, is simply given by the ratio of these time scales, as $T^\ast = k_B T / \xi_{FL}$. Two additional rescaled temperatures could be defined using $\epsilon_{FENE}$ or $\epsilon_{\text{bend}}$ as the reference energy, but we find that the most convenient description and characterization of the transition to glassy behavior is obtained using $\xi_{FL}$.

Initial loose polymer conformations are generated within a cubic periodic domain using a nonoverlapping random-walk algorithm. We use a system of $N_p = 5 \times 10^3$ beads, in chains of uniform length $L$, which we vary from 2 to 50. The value of $N_p$ is chosen following the entanglement criteria of ref 24, and moreover we demonstrate the sensitivity to $N_p$ in Figure 9, bottom left panel. For each value of $L$ we generate five realizations of the system for the purposes of ensemble averaging. We comment on the variation between realizations elsewhere.22 The system is first equilibrated in a melted state at $T^\ast = 1.2$, maintaining zero external pressure using a Nose–Hooke barostat with a damping parameter of 100. The system is subsequently cooled to $T^\ast = 0.1$ by decreasing $T^\ast$ at rate 1/\tau_c with \tau_c = 10^3 τ. Since $T^\ast = 0.1$ is below the glass transition for all of the polymers considered in this work, this cooling procedure allows us to measure $T_g$. For determining the vibrational properties, though, it is necessary to go to lower temperatures. To reach temperatures closer to $T^\ast = 0$, we subsequently relax the system further by applying the gradient method to the simulation configuration at constant volume. We used the fluctuation of net forces acting on beads $(f_i^2) / \langle f_i^2 \rangle$ as a measure for temperature. By comparing the forces with the reference value from $T^\ast = 0.1$, we get the temperature of the relaxed system configuration by $T^\ast = 0.1 \langle f_i^2 \rangle / \langle f_i^2 \rangle (T^\ast = 0.1)$. By this protocol, a target temperature of $T^\ast = 10^{-4}$ was reached for each realization. Further decreasing the temperature does not lead to changes in the VDOS or structural quantities.

III. STRUCTURE OF COARSE-GRAINED POLYMER GLASSES

III.A. Changes in $T_g$ with Chain Length and Stiffness. Ensuring that the external pressure remains zero, the system undergoes a decrease in volume $V$ as it is cooled (Figure 2). In Figure 2a, a change of gradient is identified at $T^\ast = T_g^\ast$, corresponding to the glass transition.25,26 As reported in Figure 2a,b, the model predicts that $T_g^\ast$ increases with the chain length,27 consistent with the classical free-volume result of Flory and Fox28 and with the more recent criterion based on generalized Born melting for glasses.29 This is the case for both fully flexible $(\kappa_{\text{bend}}/\kappa_{\text{LJ}} = 0)$ and very stiff $(\kappa_{\text{bend}}/\kappa_{\text{LJ}} = 17.5)$ chains. As expected,30 we further find that $T_g^\ast$ increases with $\kappa_{\text{bend}}/\kappa_{\text{LJ}}$ (Figure 2c,d), with apparent limiting values occurring for $\kappa_{\text{bend}}/\kappa_{\text{LJ}} \rightarrow 0$ and $\kappa_{\text{bend}}/\kappa_{\text{LJ}} \rightarrow 4$. The increase is significantly more pronounced as $L$ is increased. We report the increase of $T_g^\ast$ with $\kappa_{\text{bend}}/\kappa_{\text{LJ}}$ and provide further details in our earlier article.31

We find that $T_g^\ast$ varies between $\approx 0.4$ and $\approx 0.9$ for all values of $L$ and $\kappa_{\text{bend}}/\kappa_{\text{LJ}}$ and that in all cases the system is well within the glassy state at $T^\ast = 0.1$. When we increase $\kappa_{\text{bend}}/\kappa_{\text{LJ}}$ above 20 (corresponding to $\kappa_{\text{bend}}/\xi_{\text{FENE}} > 1$), we find that the angular potentials are large enough to stretch the FENE bonds beyond their maximum length $R_0$ at which point the chains break and the simulation becomes unstable due to the divergence of the $\ln(1 - (r/R_0)^2)$ term in the FENE expression. We therefore treat this as a limiting value of $\kappa_{\text{bend}}/\kappa_{\text{LJ}}$ and do not explore stiffer chains.

III.B. Changes in Density with Chain Length and Chain Stiffness. It is evident from Figure 2 that the value of $V/N_p \sigma^3$ at $T^\ast = 0.1$ is sensitive to both chain stiffness and length. In our earlier paper,5 we showed that differences in $V/N_p \sigma^3$ persist even at fixed $T^\ast / T_g^\ast$ (rather than fixed $T^\ast$), demonstrating that there are robust changes in density as chain bending and length are varied. We present in Figure 3 a contour map of polymer glass density, quantified as the number of beads per unit volume $N_p/V$, rescaled by the characteristic bead excluded volume $\sigma^3$, measured at $T^\ast = 0.1$. Since our subsequent minimization protocol conserves volume, the map also applies to $V/N_p \sigma^3$. We note that increasing chain length leads to compaction for flexible chains but expansion for stiff chains.

Figure 2. Volume–temperature curves for polymer cooling as functions of chain length and chain stiffness. The volume is rescaled with bead number $N_p$ and size $\sigma^3$. (a) Increasing chain length for $\kappa_{\text{bend}}/\kappa_{\text{LJ}} = 0$. Highlighted is the glass transition temperature $T_g^\ast$, where the polymer transitions from a melt to a glassy state. (b) Increasing chain length for $\kappa_{\text{bend}}/\kappa_{\text{LJ}} = 17.5$. (c) Increasing chain stiffness for chain length 3. (d) Increasing chain stiffness for chain length 50. Colors in (a) and (b) refer, from blue to red, to chain lengths 2, 3, 4, 5, 20, and 50. Colors in (c) and (d) refer, from blue to red, to chain stiffnesses $\kappa_{\text{bend}}/\kappa_{\text{LJ}} = 0.0175$, 0.0525, 0.175, 0.525, 1.75, 5.25, and 17.5.

Figure 3. Contour plot of polymer glass density (enumerated as $N_p \sigma^3/V$ at $k_B T / \xi_{FL} = 0.1$) as functions of chain length and chain stiffness. We note that increasing chain length leads to compaction for flexible chains but expansion for stiff chains.
at $T^* = 10^{-4}$. With respect to the density for $L = 3$, it is interesting to note that increasing chain length leads to compaction for flexible chains but expansion for stiff chains.

For very flexible chains ($\frac{\kappa_{\text{bend}}}{\kappa_{\text{LJ}}} < 1$), the key effect of increasing chain length is to move bead pairs from the LJ rest position at $2^{1/6} \sigma$ to the FENE rest position at $0.96 \sigma$, while maintaining a purely central-force system with minimal explicit bending constraints. As a result, the density increases with increasing chain length as illustrated in Figure 3 and similarly by the decreasing value of $V/N_{\sigma}^{p}$ at $T^* = 0.1$ in Figure 2a.

For less flexible chains, the roles of stiffness and chain length are more subtle. In order to achieve mechanical stability at, and below, $T_{c}^{\kappa}$, approximately monodisperse beads in central force networks require six pairwise interactions to fully constrain their translational degrees of freedom, in agreement with Maxwell’s criterion for isostaticity. As bending stiffness is increased, the translational motions of beads along chain backbones become increasingly constrained by three (and many)-body interactions. This means that as $\frac{\kappa_{\text{bend}}}{\kappa_{\text{LJ}}}$ increases, the translational degrees of freedom of individual beads can be fully constrained with fewer than six pairwise interactions per bead.\textsuperscript{29,32} We quantified this effect using the coordination number Z in another contribution.\textsuperscript{31} Since we operate at fixed external pressure, this lower coordination further implies that marginal stability can be achieved at lower density as stiffness is increased. This is the result observed in Figure 3 for $L > 3$ and indeed in Figure 2c,d (inverse density $V/N_{\sigma}^{p}$ at $T^* = 0.1$ increases with increasing $\frac{\kappa_{\text{bend}}}{\kappa_{\text{LJ}}}$, with the effect becoming more evident for longer chains, which permit many-body interactions.

As we deviate from the short chain limit, it is interesting to note that there is a continuous transition from very weak dependence to rather strong dependence on stiffness with increasing chain length. This can be interpreted by considering again the coordination argument above, which argued that bending constraints impose many-body effects along chains such that marginal stability can be achieved with fewer pairwise contacts than otherwise. For shorter chains, the maximum number of beads than can be correlated with one another in this way is small, so many-body interactions only have a weak contribution to the overall stability of the material. As such, when the stiffness is increased in short chains where there are not many angular potentials defined (in relative terms), most of the interactions remain as central force and as such the density varies only weakly. Conversely, for long chains, the increased stiffness allows many-body bending constraints to affect a higher proportion of the overall number of interactions, so the density decrease becomes exaggerated. Interestingly, at $L = 50$, the variation of density with chain stiffness is not linear, but rather it has an inflection around $\frac{\kappa_{\text{bend}}}{\kappa_{\text{LJ}}} \approx 1$.

III.C. Deviations from Rest Positions and the Resulting Internal Stresses. It is likely that steric constraints will play a role within our densely packed systems, meaning that beads will not typically be situated at their minima with respect to all three potentials ($L_{J}$, FENE, bending) even when the temperature is considerably less than $T_{c}^{\kappa}$. Such deviations from minima will lead naturally to internal stresses in the material. As chain length and bending stiffness are increased, it is likely that the extent to which beads deviate from their respective rest positions, and hence the total internal stress in the material, will change. It was shown recently\textsuperscript{33} that properly accounting for internal stresses in jammed emulsions is crucial to correctly obtaining the VDOS. Here we give a description of the source of internal stresses, which will help to guide our interpretation of the VDOS below.

To illustrate the deviation of beads from their rest positions, we present in Figure 4 an example bead–bead radial distribution function $g(r)$ and angular distribution function $p(\theta)$ for chains with $L = 10$ and $\frac{\kappa_{\text{bend}}}{\kappa_{\text{LJ}}} = 0.175$. Shown are the FENE rest position, LJ rest position, and the resting angle as well as the force cutoff beyond which we do not compute LJ interactions. (c) and (d) illustrate the implicit angular resting positions that arise due to two different configurations of the excluded volumes of beads.

![Figure 4](image)

**Figure 4.** (a) Radial distribution function $g(r)$ and (b) angular distribution function $p(\theta)$ for chains with $L = 10$ and $\frac{\kappa_{\text{bend}}}{\kappa_{\text{LJ}}} = 0.175$. Shown are the FENE rest position, LJ rest position, and the resting angle as well as the force cutoff beyond which we do not compute LJ interactions. (c) and (d) illustrate the implicit angular resting positions that arise due to two different configurations of the excluded volumes of beads.
≈ 2.5σ), which we label with the index α (so the bead–bead distance is \( r_\alpha \)). We take the average magnitude of the deviation of \( r_\alpha \) from the LJ rest position. Analogous calculations are done for FENE interactions and bending interactions. Overall, we compute \( \frac{1}{N_{\alpha}} \sum_\alpha \| r_\alpha - 2^{1/6} \sigma \| \), \( \frac{1}{N_{\text{FENE}}} \sum_\beta \| r_\beta - 0.96 \sigma \| \), and \( \frac{1}{N_{\text{bend}}} \sum_\gamma (\theta_\gamma - \theta_\gamma^{\text{rest}}) \). The results are given in Figure 5a–c for LJ, FENE, and bending interactions, respectively.

As bending stiffness \( \kappa_{\text{bend}}/\kappa_{\text{LJ}} \) increases, the chains increasingly conform to \( \theta_\gamma \), independently of chain length (Figure 5c). This requires changes in structure that must be accommodated by small additional deviations of LJ and FENE bonds from their resting positions. For LJ interactions (Figure 5a) the deviations increase steadily with stiffness. This effect is more marked for long chains, which have a larger number of bending constraints per bead. For FENE bonds (Figure 5b), meanwhile, the behavior is more subtle. The primary effect is similar to that observed for LJ interactions: increasing bending stiffness increases the deviation of FENE bonds from their rest positions. For short and flexible chains, however, there is an anomalous secondary effect: for \( \kappa_{\text{bend}}/\kappa_{\text{LJ}} \lesssim 1 \), the deviation of FENE bonds increases with decreasing chain length. As discussed above, the ubiquity of LJ interactions means they may dictate the positions of FENE bonded beads regardless of their considerably weaker potential. As chain length decreases, the average number of FENE bonds per bead decreases steadily, while the number of LJ interactions per bead remains unchanged. This means that LJ interactions can have more influence on FENE positions as chain length decreases, thus leading to additional stretching. This effect is stronger for flexible chains, which do not have the additional effect of angular constraints.

We further show that deviations from rest positions lead directly to potential energy being stored in the system in each case. The potential energy is calculated by summing \( U_{\text{LJ}}, U_{\text{FENE}}, \) and \( U_{\text{bend}} \) over every interaction in the minimized system configuration and is given as a function of chain length and stiffness in Figure 5d. The main effect is that increasing chain length introduces more FENE bonds into the system, whose individual deviations of \( \sim 0.02 \sigma \) (Figure 5b) contribute significantly to increasing stored potential energy. There is an additional effect whereby stored potential energy grows with increasing stiffness. This has contributions from LJ and FENE, in line with their deviations shown in Figures 5a,b, and also from bending potentials. Although the triplet conformations increasingly conform to their rest positions with increasing stiffness, remaining deviations become progressively more costly as \( \kappa_{\text{bend}}/\kappa_{\text{LJ}} \) increases, leading to a contribution to the stored potential energy.

### III.D. Overview of Structural Changes

The evolution of density and internal stresses (as parametrized by deviations from interaction energy minima) are strong functions of the bending stiffness and chain length. Upon increasing the stiffness, all angles between adjacent bonds tend to approach the minimum of the bending potential at \( \theta_\gamma \). However, this effect competes with the tendency of neighboring beads to stay close to the minima of LJ and FENE interactions. This competition leads to an increase of potential energy due to pairs of beads drifting subtly away from LJ and FENE minima. The effect of increasing the chain length is to insert more FENE bonds into the system. Since these are typically \( \sim 0.02 \sigma \) from their resting positions, this leads to a sharp increase in stored potential energy. We have checked that increasing chain length further above \( L = 50 \) does not bring any further evolution, and we can safely conclude that chain length has a non-negligible effect only for \( L < 50 \).

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**Figure 5.** Structural origin of internal stresses in the polymer glasses as a function of chain length and stiffness: (a) mean deviation of LJ pairs from their rest position; (b) mean deviation of FENE bonds from their rest position; (c) mean deviation of angles from their rest position, showing approximate independence of chain length; (d) total potential energy per unit volume, equivalent to the internal stress of the material.
IV. VIBRATIONAL DENSITY OF STATES

We next investigate the connection between chain length, angular potential, and the VDOS. Importantly, changes in the spectrum due to increasing the angular potential are related not only to the associated increase in angular forces but also to the structural changes that arise as discussed above. In what follows, we first outline our formalism for obtaining the VDOS, including a description of how we decompose it into various contributions. We then give an overview of the generic features of the VDOS of polymeric glasses, before focusing specifically on the behavior with respect to chain length and bending stiffness.

IV.A. Formalism for Obtaining the VDOS. Since we prepared the glasses well below $T_g^f$, we can ignore any effects of thermal noise and hence work in the harmonic approximation, where the displacements of the system around energy minima are small. The equation of motion can therefore be written with the Hessian $H$ of our system:

$$m \ddot{\mathbf{u}} = -H \mathbf{u}$$

(6)

Here $m$ is the mass of the constituent beads of our polymer chains (which we take to be uniform) and $\mathbf{u}$ is the displacement field. We can convert this equation into an eigenvalue problem by performing a Fourier transform, which gives

$$m \omega^2 \hat{\mathbf{u}} = H \hat{\mathbf{u}}$$

(7)

where $\omega$ are the eigenfrequencies of our system and $\hat{\mathbf{u}}$ are the eigenvectors (displacement fields). Arranging the eigenfrequencies in a normalized histogram gives us the vibrational density of states (VDOS) of our system. To obtain $\omega$, we first need the explicit expressions for the elements of the Hessian, after which we can solve eq 7 numerically. The elements of the Hessian are defined as second derivatives of the potential energy of the system:

$$H_{ab}^{nm} = \frac{\partial^2 U(z)}{\partial z^a \partial z^b}$$

(8a)

$$\frac{\partial^2 U(z)}{\partial z^a \partial z^b} = \frac{\partial^2 U(z)}{\partial z^a \partial z^b} + \frac{\partial U(z)}{\partial z^a} \frac{\partial U(z)}{\partial z^b}$$

(8b)

Here, $U$ represents the overall potential, consisting of the sum of $U_{lj}$, $U_{K}$, and $U_{bend}$ $z$ is a generic argument, and $a$ and $b$ label the Cartesian components. As one can see, the entries in the Hessian consist of two parts: one proportional to the spring constant $c$ between two beads and another one proportional to the tension $t$ (precise definitions are given in Appendix A). The latter contribution vanishes if all bonds are at their energy minimum at the same time. In reality, this would require perfect crystallization of the system, which is often not possible or would take a very long time. In this case, crystallization is inhibited even for fully flexible chains by the disparity in rest positions of LJ and FENE interactions.

Another source of tension is thermal noise, though this is not addressed in the present work since we work sufficiently below $T_g^f$. The main source of tension terms in our simulations is thus the angular potential and its competition with the LJ and FENE potentials, as discussed above. This combination of potentials creates two competing effects: the stronger angular potential forces all angles closer to the rest angle $\theta_0$ but also increases the strength of the tension for a given deviation, and generates additional tensions due to increased deviations from the LJ and FENE minima. Including them in the Hessian, we can now solve eq 7 and get the eigenvalues $\omega^2$ and displacement fields $\hat{\mathbf{u}}(\omega)$. The units of $\omega$ are $\sqrt{\kappa_{lj}/ma^2}$.

A particularly instructive quantity is the ratio at which different vibration patterns contribute at certain frequencies. We are especially interested in the internal-coordinate directions shown in Figure 6. These motions correspond to out-of-plane “rocking” motions (red, $\hat{a}_{i,1}$), perpendicular (to the chain) motions that remain in the plane of the chain (blue, $\hat{a}_{i,2}$), and “along-chain” motions (green, $\hat{a}_{i,3}$). To obtain each of these contributions separately, we project the displacement vector $\hat{\mathbf{u}}(\omega)$ of each bead onto the orthogonal basis formed by the three unit vectors $(\hat{a}_{i,1}, \hat{a}_{i,2}, \hat{a}_{i,3})$ (see Figure 6), generating a new representation $\hat{\mathbf{v}}(\omega)$.
is a weight function which measures the contribution of each of the three different motions discussed above, from which we get a partial VDOS showing the contribution of each of these three motions to the full VDOS. This decomposition provides insights into the dynamics of the chains at different frequencies in the spectrum.

\[ X_j(\omega) = \sum_{i=1}^{N} \tilde{v}_{ij}^2 \]

\[ j = \{1, 2, 3\} \]

**IV.B. Vibrational Spectrum: Results and Interpretation.** 1. Collective Lennard-Jones “Sea” and Higher Frequency Skeletal Modes. An example VDOS is given in Figure 6, for chain length \( L = 30 \) and stiffness \( \kappa_{\text{bend}}/\kappa_{\text{FENE}} = 0.525 \), while in Figure 9 we present the VDOS across the full range of
2. Short Chain Behavior. For chains with \( L = 3 \), we use the analytical form of the eigenvalues, including both bending and stretching interactions, to provide insights into the behavior of the VDOS measured in the simulation. The analytical derivation is reported with full details in Appendix B, with the expressions for the eigenvalues given in eq B2. We present both the analytical and numerical results in Figure 7.

The analysis predicts three nonzero eigenmodes, whose characteristic eigenfrequencies as a function of \( k_{\text{bend}}/k_{\text{FENE}} \) are given in Figure 7b. The splitting of their associated motions into along-chain and perpendicular components is given in Figure 7d,f. Since \( L = 3 \) chains are planar, rocking motions are not part of this analysis. We have seen that rocking motions contribute mostly to the lower LJ sea band and not so much to the high-frequency skeletal band which is our main focus here.

In Figure 7a,c,e we can see the VDOS for three different values of \( k_{\text{bend}}/k_{\text{FENE}} \) as determined from the simulation at \( L = 3 \). Aside from the LJ peak (which is not part of the analytical model), we can see that the VDOS follows closely the analytical prediction of the frequencies. In particular, the peak \( \omega_1 \) remains dominated by along-chain motions and also remains rather independent of \( k_{\text{bend}}/k_{\text{FENE}} \) throughout. \( \omega_2 \) is initially at zero (meaning that it is a soft mode) but becomes stiffer and moves to the right as \( k_{\text{bend}}/k_{\text{FENE}} \) is increased. Meanwhile, \( \omega_3 \) is initially slightly lower than \( \omega_1 \) but also increases progressively, eventually crossing over and becoming the higher of the three eigenvalues and at the same time becoming dominated by perpendicular motion.

While the analytical model predicts delta peaks at eigenvalues \( \omega_1, \omega_2, \) and \( \omega_3 \), in practice the peaks are broadened due to the distribution of rest angles, even for very high bending stiffness. At around \( k_{\text{bend}}/k_{\text{FENE}} \approx 0.3 \) the two peaks \( \omega_1 \) and \( \omega_2 \) start to overlap and merge into a single high peak. By looking at the evolution of eigenfrequencies and the associated motions for bending stiffness larger than \( k_{\text{bend}}/k_{\text{FENE}} = 0.3 \), we can also verify the interpretation that high-frequency modes \( \omega_3 \) are dominated by perpendicular motions for high \( k_{\text{bend}}/k_{\text{FENE}} \), whereas the mode that only depends on the backbone interaction \( \omega_1 \) is dominated by along-chain motions independently of \( k_{\text{bend}}/k_{\text{FENE}} \). The mode \( \omega_2 \) has weights for along-chain and perpendicular motion which are the specular opposite to the \( \omega_3 \) case and hence presents a growing along-chain character upon increasing \( k_{\text{bend}}/k_{\text{FENE}} \).

VDOS for additional values of \( k_{\text{bend}}/k_{\text{FENE}} \) are shown in Figure 9. We can clearly see the continuous shift of modes according to the analytical result given in eq B2, reflected also in the motion pattern associated with those modes as shown in Figure 7.

3. Dependence on Bending Stiffness for \( L > 3 \). For longer chains we can see the same general features as for \( L = 3 \) (Figures 8 and 9). Higher bending stiffness leads to redistribution of modes from the lower part of the FENE regime toward higher frequencies with an overlap happening at \( k_{\text{bend}}/k_{\text{FENE}} \approx 0.3 \), where they form a single peak. Above that value the bending interaction shifts modes associated with perpendicular motions toward higher frequencies, while the modes associated with along-chain motion stay relatively unchanged.

The strong bending interaction also causes high frequency out-of-plane motions to appear, as discussed above. The gap between the LJ sea and the FENE band is filled by modes in the same way as the third peak arises for \( L = 3 \). We can relate the peak at \( \omega = 60 \) to \( \omega_1 \) from the \( L = 3 \) model system by looking

stiffnesses and chain lengths. A common feature of this work is a distinct splitting of the VDOS into low- and high-frequency parts, particularly evident for low bending stiffness. The low-frequency part occupies the interval \( \omega = [0, \sim 40] \), while the high-frequency part extends up to \( \omega \approx 70 \) in most cases and up to \( \omega \approx 100 \) when \( k_{\text{bend}}/k_{\text{FENE}} \gtrsim 1 \). The low- and high-frequency bands are separated by a trough, whose depth and precise location in \( \omega \) are subtly dependent on \( k_{\text{bend}}/k_{\text{LJ}} \). This generic splitting of the VDOS into two bands was shown previously by Jain and de Pablo, who considered fully flexible chains only. Moreover, experimental works in polymerization have demonstrated that during periods of chain growth a single peak in the Raman intensity transforms into two peaks.\(^{35,36} \) These may be related to the two distinct bands predicted here, though a quantitative link between the Raman spectra and the VDOS reported here remains challenging and is the subject of ongoing work. We expect, furthermore, that imposing pressures greater than zero will increase the vibrational energy of both bands, thus shifting the spectrum to higher frequencies as observed experimentally.\(^{37} \) By considering the relative prefactors of the LJ and FENE potentials, we find it instructive to interpret the low-frequency part as a Lennard-Jones “sea”, that comprises weak but ubiquitous interchain LJ interactions, while the high-frequency part represents FENE bonds that are fewer in number and follow specific paths along chain backbones. Within this picture, the contributions to the VDOS coming from bending interactions are highly sensitive to \( k_{\text{bend}}/k_{\text{LJ}} \). In particular, when \( k_{\text{bend}}/k_{\text{LJ}} \) is small, we expect bending interactions to contribute frequencies comparable to, or even lower than, the LJ interactions. By contrast, when \( k_{\text{bend}}/k_{\text{FENE}} \rightarrow 1 \), we expect the bending interactions to contribute frequencies comparable to the FENE interactions. We anticipate a redistribution, therefore, of the bending contributions from the low- to high-frequency band as \( k_{\text{bend}}/k_{\text{LJ}} \) is increased.

By analyzing the motion patterns with respect to the geometry outlined in Figure 6, we can see that for flexible chains out-of-plane (rocking) motions are predominantly apparent in the low-frequency peak, while the more energetic modes mainly contain motions in the plane (skeletal vibrations). This is consistent with the low-frequency band being mainly due to LJ interactions, since they are the only interactions in the system which are not contained in the plane defined by two adjacent bonds. Consequently, the high-frequency part is mostly caused by the FENE bonds, which point along the chain backbones.

For high bending stiffness we can see a separation between along chain and perpendicular motion, of which the latter one occupies the high-frequency part. Hence, the three-body bending interaction is mostly associated with perpendicular motion, whereas the FENE interaction is rather associated with along-chain motions. This makes sense as our chains have a rest angle \( \phi_0 \approx \pi/2 \), meaning the FENE bonds point mostly along the chain direction.

For longer chains we see rocking motion arising at higher frequencies. The reason for this is that perpendicular in-plane motion caused by one triplet in the chain causes out-of-plane motion from the perspective of neighboring triplets, since they most likely do not lie in the same plane (as would be the case for a completely flat chain). As our chains are freely rotating, having a completely flat chain is very unlikely, which explains why strong bending interaction causes rocking motion at high frequencies.
and therefore the change in rigidity, as explained in previous work.\textsuperscript{19,20} This, however, does not account for the structural effect brought about by increasing stiffness, which leads to volumetric expansion and, under certain conditions (as discussed by Ness et al.\textsuperscript{31}), may lead to a decrease of shear modulus upon increasing bending stiffness, thus giving rise to a nonmonotonic dependence of the shear modulus on chain stiffness. Clearly this mechanism by which the Boson peak is changed in polymer glasses is very different from other mechanisms discussed in the literature for small-molecule or atomic glasses.\textsuperscript{39,40}

To summarize, we can separate the spectrum into four distinct parts: (i) the lowest frequency band of LJ sea which gets lowered as, with higher bending stiffness, more modes are shifted to higher frequencies; (ii) a stationary peak around $\omega = 60$ strongly associated with along-chain motion/vibration and with characteristic frequency $\omega_1$; (iii) modes associated with vibrations perpendicular to the chain axis, which resemble the behavior of $\omega_2$ and the frequencies of which diverge with the bending stiffness; and (iv) modes associated with along-chain vibrations that resemble the behavior of $\omega_3$ filling the regime between the LJ sea and the $\omega_1$ peak.

4. Dependence on Length $L$. By increasing the chain length, we introduce more high-energy FENE bonds into the system, which leads to a shift or redistribution of modes from the low to the high-frequency band, causing a lowering of the LJ peak (Figure 9). This is particularly evident for fully flexible chains. Additionally, the number of possible polymer conformations increases drastically with chain length. In the VDOS this leads to a loss of distinct features (i.e., sharp peaks become broadened), especially in the high-frequency band. The effect is most visible for fully flexible short chains of $L = 3, 4, 5$, whereas the difference between $L = 10$ and $L = 50$ is only marginal. The total number of additional FENE bonds per chain $n$ decreases as the chains become longer $n/N = \frac{L - 1}{L}$, $\Delta n/N = \frac{1}{(L + 1)^2}$, and therefore the change in total bond energy becomes smaller. As such, the averaged spectrum of sufficiently long ($L > 5$) chains already approximates the spectrum of an infinitely long chain quite well. A sample of chains with $L = 10$ consequently shows the same shape as for $L = 50$ (or even $L = 1000$). We thus limited our analysis to systems with those lengths, as there is no new physics to see in the spectrum of chains longer than $L = 50$. The distributions of motion patterns do not change much with the chain length, aside from the adjusting to the overall shape of the spectrum described above. An analytical derivation for fully flexible linear chains with $L = 2, 3, 4, 5$, with stretching interactions, is reported with full details in Appendix C.

V. CONCLUDING REMARKS

We presented a systematic analysis and interpretation of the structure, internal stresses, and vibrational spectra of glassy polymers from coarse-grained simulations, based on the Kremer–Grest bead–spring model with an energy minimum for angular bending interaction. Varying the angular stiffness and the chain length leads to rich phenomenology: an increase in both of these parameters causes a buildup of internal stresses due to the competition between bending and stretching degrees of freedom, both of which want to minimize their energy at the same time. This leads to increased deviations from the minima of LJ and FENE interactions, an effect amplified for longer chains.
For flexible chains with $\kappa_{\text{bend}}/\kappa_{\text{FENE}} < 0.3$ there are two bands in the VDOS spectra, corresponding to LJ-dominated interactions at low frequency (the LJ sea) and to skeletal modes dominated by FENE bonds at high frequency (the high-frequency skeletal band). For chains with higher bending stiffness this separation breaks down as modes associated with angular interactions appear, filling the gap between the two bands and creating additional high frequency modes. The latter of these are mostly made up from vibrations perpendicular to the chain, while the gap is filled by along-chain motions, creating a new separation between vibrational regimes in the spectrum.

For short chains, the spectra feature sharp peaks whose behavior as a function of stiffness correspond very well with our analytical prediction. For longer chains this structure loses its sharp features and tends to a more continuous spectrum in which the various peaks are broadened by the averaging over many different chain conformations.

This framework and these concepts can be applied in future work to molecular and atomistic simulations of realistic materials, possibly in combination with the Kernel Polynomial Method which can greatly speed up the evaluation of the VDOS using the Hessian as input. We anticipate that the generic features of the VDOS predicted in this work will be robust to the introduction of more specific chemical interactions (including those that break the isotropic interaction symmetry), since the features of the vibrational spectrum are related essentially to the energy and the relative strength of interactions. For example, hydrogen bonds (which have typical energy (4–13 kJ/mol) an order of magnitude less than covalent C–C bonds (346 kJ/mol)) would be expected to add to the part of the spectrum that is already dominated by the LJ sea. The same can be stated about stacking interactions, which have typical energies of 8–12 kJ/mol.

Finally, our results may open up the possibility of quantitatively linking the Raman and Brillouin spectra of glassy polymers with their viscoelastic response, since the VDOS is a key input to calculate viscoelastic moduli within recent developments in the nonaffine linear response of amorphous solids.
**APPENDIX A. EXPLICIT FORM OF THE HESSIAN**

Here we show the explicit form of the entries to the Hessian for each potential. First we recall the general form

\[
H_{nm}^{ab} = \frac{\partial^2 U(r)}{\partial r^a_n \partial r^b_m},
\]

where \(a\) and \(b\) label the Cartesian components, and the entries are given, for a generic argument \(z_i\), as

\[
\frac{\partial^2 U(z)}{\partial y \partial y} = \frac{\partial^2 U(z)}{\partial z \partial z} \frac{\partial z}{\partial y} + \frac{\partial U(z)}{\partial z} \frac{\partial^2 z}{\partial y \partial y}.
\]

For the two central-force potentials (FENE and Lennard-Jones) we have \(z = |r_i - r_j| = r_{ij}\). It should be noted that all derivatives are evaluated at the actual system conformation from the simulation. For central forces we get

\[
H_{nm}^{ab} = \frac{\partial^2 U(r_{ij})}{\partial r^a_n \partial r^b_m} = c_{ij} \frac{\partial^2 U(r_{ij})}{\partial r^a_n \partial r^b_m} + t_{ij} \frac{\partial^2 r_{ij}}{\partial r^a_n \partial r^b_m},
\]

\[
c_{ij} = \frac{\partial^2 U(r_{ij})}{\partial r^a_n \partial r^b_m}, \quad t_{ij} = \frac{\partial^2 r_{ij}}{\partial r^a_n \partial r^b_m},
\]

\[
\frac{\partial^2 r_{ij}}{\partial r^a_n \partial r^b_m} = \frac{r_{ij}}{r_{ij}} (\delta_{ij} - \bar{n}_{ij}) \bar{n}^a_n \bar{n}^b_m.
\]

(A2)

Here \(\bar{n}_{ij} = r_i / r_{ij}\) denotes the unit bond vector between beads \(i\) and \(j\). The above expressions are valid both for FENE and Lennard-Jones bonds with the only difference being the stiffness \(c_{ij}\) and tension \(t_{ij}\) that have to be evaluated depending on the potential. For the angular potential we have a slightly different situation as our variable is now the angle between the two bonds \(r_i - r_j\) and \(r_k - r_l\):

\[
z = \theta_{ij} = \arccos \left( \frac{(r_i - r_j) \cdot (r_k - r_l)}{|r_i - r_j||r_k - r_l|} \right) = \arccos A_{ij}
\]

(A3)

To make the calculation easier, we rewrite the first line of eq A2 to give

\[
\frac{\partial^2 A}{\partial r^a_n \partial r^b_m} = \frac{1}{r_{ij}} (\delta_{ij} - \bar{n}_{ij}) (\delta_{nm} - \bar{n}_{nm})(1 - \bar{n}^a_n \bar{n}^b_m)
\]

(A4)

These formulae were derived in a slightly different but substantially equivalent fashion by Van Workum et al.\(^{14}\)

**APPENDIX B. ANALYTICAL SOLUTION FOR TRIATOMIC MOLECULE WITH BENDING STIFFNESS**

We next write down the Hessian for an isolated oligomer with \(L = 3\) (a triatomic molecule model), accounting for both stretching and bond-bending interactions. As the eigenvalues of the Hessian are invariant under spatial rotations, we can choose the chain lying flat in the \(x-y\) plane with beads \(P_1 = -r(\zeta, 0), P_2 = r(0, \nu), P_3 = r(\zeta, \theta)\) and \(\zeta = \sin \theta / 2, \nu = \cos \theta / 2\):

\[
H = \frac{\kappa}{m} \begin{pmatrix}
\zeta + \gamma' \nu^2 & -(1 - \gamma') \zeta \nu & -\zeta^2 & (1 - 2\gamma') \zeta \nu & -\gamma' \zeta^2 & \gamma' \zeta \\
-(1 - \gamma') \zeta \nu & \zeta^2 + \gamma' \zeta^2 & \zeta \nu & \gamma' \nu & \nu^2 & \gamma' \nu \\
-\gamma' \nu^2 & \gamma' \nu & -\zeta^2 & -\zeta^2 & -\zeta^2 & \nu^2 \\
(1 - 2\gamma') \zeta \nu & -\nu^2 - 2\gamma' \zeta^2 & 0 & 2\nu^2 + 4\gamma' \zeta^2 & -(1 - 2\gamma') \zeta \nu & -\nu^2 - 2\gamma' \zeta^2 \\
-\gamma' \nu^2 & \gamma' \nu & -\zeta^2 & -\zeta^2 & -\zeta^2 & (1 - \gamma') \zeta \nu \\
\gamma' \zeta & \gamma' \nu & \nu^2 & -\nu^2 - 2\gamma' \zeta^2 & (1 - \gamma') \zeta \nu & \nu^2 + \gamma' \nu^2
\end{pmatrix}
\]

(B1)

Here we used the dimensionless bending stiffness \(\gamma' = \gamma/(kr^2)\) for bond length \(r\), where \(\gamma\) (which has units of energy) is the second derivative of the angular bending potential with respect to the angle, while \(\kappa\) is the spring constant of the bond for central-force stretching of the bond. The above matrix has three nonzero eigenvalues, leading to the following eigenfrequencies:

\[
\omega_1^2 = \frac{\kappa}{m} (2 - \cos \theta)
\]

(B2a)

\[
\omega_2^2 = \frac{\kappa}{2m} \left( 2(1 + 2\gamma') + (1 - 2\gamma') \cos \theta \right) - \sqrt{\left( 2(1 + 2\gamma') + (1 - 2\gamma') \cos \theta \right)^2 - 24\gamma'}
\]

(B2b)

\[
\omega_3^2 = \frac{\kappa}{2m} \left( 2(1 + 2\gamma') + (1 - 2\gamma') \cos \theta \right) + \sqrt{\left( 2(1 + 2\gamma') + (1 - 2\gamma') \cos \theta \right)^2 - 24\gamma'}
\]

(B2c)
As we can see, one eigenfrequency $\omega_1$ is independent of the bending stiffness, while $\omega_2$ shows a convergent behavior against $\gamma'$, $\omega_2 \to \sqrt{3x/(m(2 - \cos \theta))}$ and $\omega_3$ diverges like $\sim \sqrt{\gamma'}$.

## Appendix C. Analytical Solutions for Fully Flexible Chains

In this Appendix we consider a toy model for the determination of the skeletal vibration modes of a single polymer chain. The following assumptions are made: (i) the chain is fully flexible (vanishing angular stiffness); (ii) only in-plane motions are considered (rocking or other out-of-plane vibrations are neglected). These assumptions are needed to obtain analytical results. We will start with the simplest case of a zigzag regular chain with a single fixed value $\theta$ of the angle between two adjacent bonds, and we will subsequently consider the case of a distributed $\theta$. We consider two variations of this model, first for a zigzag chain with fixed value of the angle, and subsequently for a uniform (random) distribution of the angle.

The Hessian has the following block structure:

$$
H = \begin{pmatrix}
A_{12} & -A_{12} & 0 & 0 & 0 & 0 \\
-A_{12} & A_{12} + A_{23} & -A_{23} & 0 & 0 & \cdots \\
0 & -A_{23} & A_{23} + A_{34} & -A_{34} & 0 & 0 \\
0 & 0 & -A_{34} & A_{34} + A_{45} & -A_{45} & \ddots \\
\vdots & \vdots & \vdots & \vdots & \ddots & \vdots \\
0 & 0 & 0 & 0 & \cdots & -A_{n-1, n-1}
\end{pmatrix}
$$

(C1)

with blocks given by

$$
A_{ij} = \frac{k}{m} \begin{pmatrix}
n_i n_i^{*} & n_i n_i^{*} & n_i n_i^{*} & n_i n_i^{*} \\
n_i^{*} n_i^{*} & n_i^{*} n_i^{*} & n_i^{*} n_i^{*} & n_i^{*} n_i^{*}.\end{pmatrix}
$$

(C2)

where the unit vector $n_i$ which goes from bead $i$ to a nearest-neighbor $j$, and spring constant $k = 1$. To get the characteristic polynomial $p(\lambda)$, one has to evaluate the determinant $|H - \lambda I|$. We can iteratively solve this by using the standard formula for block matrices:

$$
\begin{vmatrix}
A & B \\
C & D
\end{vmatrix} = |D||A - BD^{-1}C|
$$

(C3)

where the entries are matrices and the relation for our $3 \times 3$ blocks:

$$
A \cdot B \cdot A = \frac{k^2}{m^2} \cos^2 \theta_{AB} A
$$

(C4)

where $\theta_{AB}$ is the angle between the bonds belonging to $A$ and $B$. After some calculation we get the following recursion formula (omitting the $2L + 1$ trivial eigenvalues $\lambda = 0$) for the above matrix:

$$
p_1(x) = 1, \\
p_1(x) = x, \\
p_n(x) = x p_{n-1}(x) - \cos^2 \theta_{n-1,n-2} p_{n-2}(x)
$$

(C5)

where $x = \frac{m}{k} \lambda^2 - 2$. The recursive relation for $p_n(x)$ can also be derived on more formal grounds.$^{46}$

Note that $n$ denotes the number of bonds in a chain and not the number of beads in the chain ($n = L - 1$). For arbitrary angles between the bonds it is not possible to describe the roots of this polynomial, except for oligomers (see below). But, if all angles are the same, we can bring (C5) into the form of the Chebyshev polynomials of the second kind $U_n(x)$ by substituting $\tilde{x} = x/\cos \theta$:

$$
\tilde{p}_n(\tilde{x}) = \cos^2 \theta U_n(\tilde{x}), \\
\tilde{x} = \frac{x}{2 \cos \theta} = \frac{\frac{m}{k} \lambda^2 - 2}{2 \cos \theta}
$$

(C6)

The roots of $U_n(x)$ are $x_k = \cos\left(\frac{k}{n+1}\pi\right)$; $k = 1, \ldots, n$, which gives us the eigenvalues of the linear chain with constant angle as

$$
\lambda_k = \omega_k^2 = 2k \left[1 + \cos \theta \cos \left(\frac{kr}{n+1}\right)\right] = \frac{2k}{m}\left[2 \cos \theta - \left(\omega^2 - \frac{2k}{m}\right)\right],
$$

$$
D(\omega) = \frac{2}{\pi} \sqrt{\frac{m}{k} \cos^2 \theta - \left(\omega^2 - \frac{2k}{m}\right)^2}, \quad (\theta \neq \pi/2),
$$

$$
\rho(\omega) = \delta \left(\omega - \frac{2k}{m}\right), \quad (\theta = \pi/2)
$$

(C7)

This result can also be derived using a different approach which exploits the periodicity of the chain with constant angle, as was done by Kirkwood.$^{12}$. If we assume that the chain points along the $x$-axis we can identify the previously introduced along-chain and perpendicular motion as $A$ and $B$ in eq 6 of Kirkwood.$^{12}$ By using the dispersion relation found in this work, we can solve for those two quantities and find the weight functions:

$$
X_A(\omega) = \frac{|A|^2}{|A|^2 + |B|^2} = \frac{\cos \theta + \omega^2 + \frac{2k}{m}(\cos \theta - 1)}{2\omega^2},
$$

$$
X_i(\omega) = \frac{|B|^2}{|A|^2 + |B|^2} = \frac{\cos \theta - 1 - \omega^2 - \frac{2k}{m}(\cos \theta + 1)}{2\omega^2}
$$

(C8)

As mentioned before, for the flexible case with distributed angles an analytical solution is not accessible for a chain of arbitrary length. But we can give the eigenvalues in the case of short chains with $L = 2, 3, 4, 5$:
applications of raman spectroscopy.

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Notes

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