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Simulation study of pressure and temperature dependence of the negative thermal expansion in Zn(CN)$_2$

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(Dated: January 21, 2014)

Pressure and temperature dependence of the negative thermal expansion in Zn(CN)$_2$ is fully investigated using molecular dynamics simulations with a built potential model. The advantage of this study allows us to reproduce the exotic behaviours of the material, including the negative thermal expansion (NTE), the reduction of NTE with elevated temperature, the pressure enhancement of NTE and the pressure-induced softening. Results of the study provide us detailed data to link the properties in the energy space and the real space, giving us insights to understand the properties and the connections between them.

PACS numbers: 65.40.-b, 65.40.De, 63.20.Dj, 02.40.-k

I. INTRODUCTION

Negative thermal expansion (NTE) is a rare and counter-intuitive phenomenon found primarily in low density materials with crystal structures that are networks of linked coordination polyhedra. The study of these materials is not only of fundamental scientific importance, but also has many technological applications such as aerospace technology, optics, and electronics. While much effort has been put into finding new materials and investigating the origin of NTE in them, much less attention has been paid to the change in NTE behaviour subject to heating and stress, which holds great importance for the possible applications of the material. For example, due to stresses and heating, problems such as phase transitions of the NTE filler and thermal expansion mismatch between the NTE filler and matrix are always encountered in designed composites with tailored thermal expansion.

In this paper, we conduct a simulation study of Zn(CN)$_2$ focusing on the pressure and temperature effects on its negative thermal expansion. We chose this material for several reasons. Firstly, Zn(CN)$_2$ is a well-known representative NTE material. It has a framework structure consisting of tetrahedral groups of atoms linked by diatomic rods of C–N and has exceptionally large isotropic NTE of $\alpha_{\text{linear}} = -16.9$ MK$^{-1}$ (twice as large as that of ZrW$_2$O$_8$). Secondly, the material shows a variety of exotic properties in experiments including reduction of its NTE on heating, pressure-enhanced thermal contraction, and pressure-induced softening, none of which are fully understood. Thirdly, with previous DFT calculations of Zn(CN)$_2$ explaining the origin of NTE of the material in terms of Grüneisen theory, it would be useful to draw a clear link between the values of the Grüneisen parameters in energy space and the structural vibrations in real space with full anharmonicity (which should be important in such an NTE system) based on theoretically reproducing the aforementioned exotic properties.

Here, we have built a Zn(CN)$_2$ potential model based on first-principles calculations. Lattice-dynamic calculations and large-scale molecular dynamics (MD) simulations were carried out for the material using this model which was justified by comparing against the available experimental data. The results in both energy and real space provide us fundamental clues to understand the NTE as well as the related exotic behaviours of Zn(CN)$_2$.

II. BUILDING THE MODEL

We started with total-energy calculations for a [Zn(CN)$_4$]$^{2-}$ cluster. The $-2$ charge comes from the fact that every zinc is shared by four neighbour atoms (C or N), and that the bond should be highly ionic according to an initial judgment of zinc having much smaller electronegativity than carbon and nitrogen. Four CN$^-$ molecular ligands around each Zn$^{2+}$ would give us a $-2$ charge on the cluster. Geometry optimization of the clus-
TABLE I. The potential model. E is energy, r is inter-atomic distance, r_0 is equilibrium inter atomic distance, \( \theta \) and \( \varphi \) are bond angles, and \( \theta_0 \) is equilibrium bond angle. Energies are in eV, distances are in Å and bond angles are in degrees. The prefactor of the harmonic bond-bending term is in eV/\( \text{rad} \).

| Potential | Form of the Potential | Type of Bond | Values of Parameters |
|-----------|-----------------------|--------------|----------------------|
| Morse potential | \( E_M = D \left[ (1 - \exp (-\alpha (r - r_0)))^2 - 1 \right] \) | Zn–C | \( D = 0.2432, \alpha = 2.917, r_0 = 2.109 \) |
| | | Zn–N | \( D = 0.1795, \alpha = 2.701, r_0 = 2.157 \) |
| Harmonic three-body potential | \( E_H = (1/2)K(\theta - \theta_0)^2 \) | C–Zn–C | \( K = 1.5157, \theta_0 = 109.47 \) |
| | | C–Zn–N | \( K = 1.2695, \theta_0 = 109.47 \) |
| | | N–Zn–N | \( K = 1.0233, \theta_0 = 109.47 \) |
| Linear three-body potential | \( E_L = K(1 - \cos \varphi) \) | Zn–C–N | \( K = 1.1968 \) |
| | | Zn–N–C | \( K = 0.6359 \) |
| Buckingham potential | \( E_B = A \exp(-r/\rho) - (C/r^6) \) | C–C | \( A = 2806.9, \rho = 0.2667, C = 17.67 \) |
| | | C–N | \( A = 2365.0, \rho = 0.2825, C = 20.88 \) |
| | | N–N | \( A = 1992.7, \rho = 0.2874, C = 24.67 \) |

...ter resulted in a perfect tetrahedral conformation.

Total energies for different configurations of the cluster (with bond stretching and angular distortions) were computed using DFT in GAMESS(US) with the PBE0 functional.\(^{14,15}\) Correlation-consistent basis sets up to aug-cc-pVQZ were tested, and an aug-cc-pVTZ basis set was found to have sufficient accuracy and no significant basis set superposition errors (BSSE) at different configurations.

Various interatomic potential forms were then fitted to the calculated energy curves to obtain the initial potential parameters. The short-range interactions are the Morse potential for Zn–C/N, a harmonic three-body-bond-bending term for C/N–Zn–N/C and a linear-three-body term for the angular distortion of Zn–C/N–N/C. The long-range van der Waals interactions are described by a Buckingham potential with parameters from Williams.\(^{16}\)

Two types of clusters with Zn–C–N order and Zn–N–C order are used. The multipole on each cluster was calculated by distributed multipole analysis (DMA)\(^{17}\) using CamCASP\(^{18}\). The effective point charges on the atoms were then obtained by fitting to the electrostatic potential from the rank 4 (hexadecapole) distributed multipoles using the MULFIT program.\(^{19,20}\) Fig. 1 shows the difference in the electrostatic potential between the point-charge model and the DMA result. The root mean square of the difference is less than 4 kJ/mol and 8 kJ/mol for clusters of Zn–C–N and Zn–N–C, respectively, corresponding to about 1% relative difference in the electrostatic potential around the clusters. The averaged effective point charges on each atom are (in electron units) +1.14 for Zn, −0.21 for C and −0.36 for N.

The initial potential model was then refined by refitting to DFT energy surfaces of various cluster configurations. The point charges were fixed during this process.

This gave us final potential parameters that implicitly incorporate the effects of higher-ranking multipole moments and atomic polarization. Due to the high strength of the C–N bond,\(^{21,22}\) this group was treated as a rigid rod in all cases. The potentials with their parameters are listed in Table I.

III. COMPUTATIONAL METHODS

Harmonic lattice dynamics (HLD) and quasi-harmonic lattice dynamics (QHLD) calculations were carried out using GULP.\(^{23}\)

Molecular dynamics (MD) simulations were performed using DL_POLY\(^{24}\) for a 10 x 10 x 10 supercell containing 10000 atoms with periodic-boundary conditions. A constant stress constant temperature (NVT) ensemble with a Nosé-Hoover thermostat\(^{25}\) was used. The long-range Coulomb interactions were calculated using the Ewald method with precision of 10\(^{-6}\). The equations of motion were integrated using the leapfrog algorithm with a time step of 0.001 ps. A total of 20000 time steps were used to achieve equilibration. At different temperatures and pressures, snapshots of atomic trajectories after equilibration were recorded every 0.02 ps up to a total of 50 ps for the follow-up analysis. Both ordered model with \( Pm3m \) symmetry and disordered model with \( Pm3m \) were used. The latter was constructed by randomly switching C and N atoms in the supercell.

IV. AN INITIAL TEST OF THE MODEL

We compared some basic quantities obtained from the model against experiment. The optimized structure gave the cell parameter as \( a = 5.9176 \) Å, and the Zn–C/N bond length as \( r = 1.9782 \) Å compared to \( a = 5.9227(1) \) Å and \( r = 1.9697(3) \) Å from an experiment at 14 K.\(^{22}\) The model is found to be stable in lattice-dynamic calculations, and the calculated phonons are in good agreement with the spectroscopy data,\(^{21}\) as shown in Table I. The dispersion curves shown in Fig. 2 generally agree well with previous DFT calculations.\(^{20,21}\)
TABLE II. Calculated phonon frequencies at the Γ point for both CN ordered and CN disordered (virtual crystal) models. Experimental Infra-red and Raman data and DFT calculated results are provided for comparison. Frequencies are in cm$^{-1}$.

| Mode     | $T_{1u}$ | $T_{2g}$ | $E_{g}$ | $T_{2g}$ | $T_{1u}$ |
|----------|----------|----------|---------|----------|----------|
| Ordered  | 173      | 186      | 328     | 337      | 463      |
| Disordered | 210     | 220      | 314     | 349      | 490      |
| Experiment | 178     | 216      | 334     | 339      | 461      |
| DFT$^{13}$ | 178     | 204      | 330     | 336      | 476      |

FIG. 2. Zn(CN)$_2$ dispersion curves calculated along high-symmetry directions in the Brillouin zone. Black curves are the phonons of the zero-pressure cell; light-red curves are the phonons of the cell at 1.0 GPa hydrostatic pressure. The acoustic modes at M (0.5, 0.5, 0.0) and at the midpoint of Γ–R (0.25, 0.25, 0.25) are the first to become unstable at high pressure.

It is worth noting the close similarity between the dispersion relations of wave vectors along Γ–X–R and R–M–Γ, as well as the near-mirror symmetry of the dispersion curves along Γ–R (the middle line of Γ–R is the mirror line). This makes sense given the existence of the two interpenetrating cristoablite-like networks in the material. For the lowest-frequency mode at wave vector R, the two networks would translate like acoustic modes but out of phase with each other, leading to a dispersion relation that has the appearance of an acoustic mode and a positive Gr"{u}neisen parameter but with a non-zero band gap. The frequency of this mode is 0.16 THz, lower than the values of 0.69 THz and 0.48 THz from the DFT calculations in Ref. [10] and Ref. [13] respectively, suggesting a softer long-range interaction between the two networks in our model.

FIG. 3. Zn(CN)$_2$ isotherms calculated for pressures 0.0–0.7 GPa in 0.1 GPa increments. Red lines are calculated values; black markers are from neutron diffraction data. The inset shows the enhancement of NTE under pressure, observed in both calculations and in experiment.

FIG. 4. NTE in Zn(CN)$_2$ calculated using MD. Symbols are data from the CN-disordered system, solid lines are data from the CN-ordered system. CN ordering thus has a negligible effect on the NTE behaviour.

V. THERMODYNAMIC PROPERTIES

A. Pressure and temperature dependence of the NTE

The calculated NTE curves of Zn(CN)$_2$ under different pressures from 0.0 to 0.7 GPa with increments of 0.1 GPa are shown in Fig. 3. The pressure-enhanced α (averaged over 50–300 K) at 0.0, 0.2 and 0.4 GPa are $-12.62$, $-14.09$ and $-15.97$ MK$^{-1}$, respectively, compared to the experimental values of $-17.40(18)$, $-18.39(27)$ and $-19.42(23)$ MK$^{-1}$ at the corresponding pressures. The MD successfully captured the gradual reduction of NTE on heating which has been observed in X-ray scattering.
According to the MD, $\alpha_V$ is $-30.3$ MK$^{-1}$ at 300 K, much lower than the value at 25 K, $-47.3$ MK$^{-1}$.

Experiment$^{23}$ has confirmed that Zn(CN)$_2$ exists in a disordered form with $Pn\bar{3}m$ symmetry, i.e. the carbon and nitrogen atoms are randomly placed on their symmetric sites in the structure, compared to the ordered model with $P\bar{3}m$ symmetry. NTE of the disordered model from the MD calculation is displayed as symbol plots in Fig. 4 compared to that of the ordered model in solid lines. Clearly, there is no significant difference between the NTE of these two systems.

B. Pressure and temperature dependence of the mechanical properties

The bulk modulus $B$ is directly related to the coefficient of thermal expansion $\alpha_V$ according to the classical form

$$\alpha_V = \frac{\pi c_v}{B}$$

where $c_v = \sum \frac{\hbar \omega_s/V}{\pi} \left( \frac{\partial n_s}{\partial T} \right)$ is the specific heat and the sum is over all modes with frequencies $\{\omega_s\}$. $n_s = [\exp(\hbar \omega_s/k_B T) - 1]^{-1}$ is the Bose-Einstein relation with $k_B$ the Boltzmann constant. The overall Grüneisen parameter $\bar{\gamma}$ is calculated by summing over all mode Grüneisen parameters weighed with their contribution to the specific heat.

At 300 K, the values of the bulk modulus $B_0$ and its first derivative with respect to pressure $B'_0$ at zero pressure obtained by using the 3rd-order Birch-Murnaghan equation of states to fit to the isotherms calculated from our MD are $B_0 = 34.47(31)$ GPa and $B'_0 = -4.2(5)$. Both are in good agreement with the experimental values of 34.19(21) GPa and $-6.0(7)$, respectively$^{10}$. However, the values of the bulk modulus obtained from the previous DFT calculation$^{10,13}$ are much higher than the experiment.

We obtained the bulk modulus of Zn(CN)$_2$ at different temperatures and pressures by numerically computing derivatives of the $p-V$ data from the MD simulations. As shown in Fig. 5 on cooling from 300 K to 50 K, $B$ increases by $26\%$ compared to the experiment of $15\%^{13}$. Note that $B$ doesn’t change much with pressure, but changes largely with temperature. Referring to Fig. 3 we found that a $5\%$ decrease in volume caused by heating at zero pressure corresponds to as much as $60\%$ decrease in the bulk modulus, while the same amount of volume decrease caused by compression would only reduce the bulk modulus by less than $5\%$. This means that the bulk modulus of the material not only depends on the volume change per se, but also on the means of changing the volume — by heating or compression. This breaks Birch’s law of corresponding states$^{20,22}$. The same anomaly has been observed experimentally in ZrW$_2$O$_8$, where the bulk modulus decreases by $40\%$ on cooling from 300 K to 0 K$^{23}$.

According to the thermodynamic expressions of $\alpha_V = \partial(\ln V)/\partial T$ and $B = \partial p/\partial(\ln V)$ combined with Maxwell relation $\partial^2 V/(\partial T\partial p)_{p,T} = \partial^2 V/(\partial p\partial T)_{T,p}$, the pressure enhanced NTE of the material follows naturally from the relation

$$\left( \frac{\partial \alpha_V}{\partial p} \right)_T = \frac{1}{B^2} \left( \frac{\partial B}{\partial T} \right)_p .$$

FIG. 5. Pressure and temperature dependence of the bulk modulus of Zn(CN)$_2$. Results calculated using the $p-V$ data from MD simulations and standard thermodynamic relations. Pressure has far less influence on the bulk modulus than temperature.

FIG. 6. Pressure and temperature-dependence of Zn(CN)$_2$ volume, calculated using MD. The inset shows the change in volume with pressure up to 3.0 GPa at 300 K. A high-pressure phase settles down after the clear discontinuity at 2.1 GPa.
Since the temperature dependence of $B$ is negative as shown in Fig. 5, $\alpha_V$ would become more negative on compression. This is consistent with what seen in the positive expansion materials where $\alpha_V > 0$ so that $\alpha_V$ would decrease on compression.

We found that the value of $B_0'$ from a lattice-dynamic calculation using GULP is 7.2 compared to its negative value at 300 K. This suggests that all the mechanical contributions at $T = 0$ are from the Zn–C/N bonds that become stiffened on compression. With elevated temperature, one can imagine that the Zn–C/N–N/C angle flexing starts to contribute to the change of volume on pressure. This mechanism costs much less energy than compressing the Zn–C/N bonds as suggested by values of the parameters in the Morse potential and the linear-three-body potential of the model (Table I). As a result, the bulk modulus decreases on compression hence the negative $B_0'$, i.e. pressure-induced softening of the material. At high temperature, $B_0'$ is expected to become less negative on heating due to the rising energy cost of further increasing the Zn–C/N–N/C angle vibrational amplitude on compression. The temperature dependence of $B_0'$ is specially discussed in our other work.\(^{29}\)

VI. PHASE TRANSITIONS

In Fig. 6 simulations were conducted for Zn(CN)\(_2\) at different temperatures in a much broader range of pressure from $-2.0$ to $3.0$ GPa, and there is clearly a phase transition of the material caused by compression at each temperature. With elevated temperature, the phase-transition pressure indicated by the discontinuity in the volume change increases, implying that the phase transition may be triggered by some soft modes that can be stabilized on heating due to the anharmonic term in their frequencies. Indeed, as shown in Fig. 2, if we compare the dispersion curves calculated at a pressure beyond 1.0 GPa (in light red) to that at zero pressure (in black), we find softening of the acoustic modes around the zone boundaries, especially the modes at M and the mid-point of $\Gamma$–R (0.25, 0.25, 0.25) which are the first ones to become unstable. The concurrent softening of the optic modes ($\sim 1.5$ THz) directly above these acoustic modes suggests a possible hybridization between the acoustic modes and the optic modes, resulting in a $k^2$ energy behavior at $k \neq 0$. Same result was found for the disordered model.

The inset of Fig. 6 shows the change of volume with pressure up to 3.0 GPa at 300 K. The volume discontinuities at 1.2 and 2.1 GPa may suggest hysteresis in the phase transition. We found that the new high-pressure phase is orthorhombic with $P2_12_12_1$ space group ($a = 10.72$ Å, $b = 10.78$ Å, $c = 10.88$ Å). This is compared to an orthorhombic phase with $Pmc2_1$ space group found beyond 1.3 GPa in the X-ray diffraction experiment.\(^{25}\)

VII. PHONONS

A. Density of states

The Fourier transformation of the atomic velocity auto-correlation function (VACF) gives us the phonon density of states (DoS) of the material. To obtain the DoS at different pressures and temperatures, we used the trajectory data of atoms from MD to calculate the VACF of Zn(CN)\(_2\).

The correlation function $C(t)$ at time $t = n\Delta t$ can be expressed as,

$$C(t) = \frac{1}{N(M - n)} \sum_{j=1}^{N} \sum_{m=1}^{M-n} v_j(m\Delta t)v_j(m\Delta t + t) \quad (3)$$

where $v_j$ is the velocity component of the $j$th atom. $\Delta t$ is the time interval of 0.02 ps. $M$ is the total number of time steps. The system was simulated for a total of 50 ps which corresponds to $M = 2500$. The correlation function was calculated for each atom with a time length of 30 ps ($n = 1500$). A Gaussian profile was used before Fourier transformation to suppress the ripple effect caused by time cut-off. The VACF of angular velocities of the C–N rigid rods rotating about their center of mass was also calculated. The Fourier transformation then gave us the DoS of the pure rotational modes of these rigid rods. To obtain the phonon spectrum, the calculated DoS was first multiplied by a weighting factor $4\pi b_k/m_k$ containing the scattering length $b_k$ and the atomic mass $m_k$ of the $k$th atom. Then, in order to mimic experimental resolution, the DoS is convolved with a Gaussian with FWHM of 10% of the energy transfer. Fig. 7 shows the good agreement between the calculated spectrum and the experiment.\(^{25}\)
In order to draw links between the atomic vibrations and the phonon properties, we computed both the overall DoS and the angular DoS of rigid rod C–N, as shown in Fig. 8 and 9. Fig. 8(a) and (b) shows the DoS at ambient temperature (300 K) under different pressures for the disordered and ordered model, respectively. The acoustic peak around 0.5 THz is softened on compression. The peaks at around 14 THz, corresponding to pure Zn–C(N) bond flexing, broaden and decrease in frequency on heating.

Previously, the peak around 0.5 THz has been found to be the major contributor to the NTE in both experiment\cite{9} and calculation\cite{10,11} due to its large negative $\gamma_{k,\lambda}$. However, if one can further identify the corresponding real-space picture of the vibrations, the reason of why the peak has the most negative $\gamma_{k,\lambda}$ compared to the other optic peaks can be revealed.

To understand the nature of various peaks in the DoS, we decided to categorize the vibrational modes in the material using the rigid unit mode model\cite{12}. First, we calculated $\gamma_{k,\lambda}$ from phonon frequencies of expanded and contracted ($\pm$0.01%) unit-cell volumes. We then coloured the dispersion curves according to both magnitudes and signs of $\gamma_{k,\lambda}$, as shown in Fig. 10(a). This representation highlights the most important phonon branches responsible for NTE of the material, namely the low-lying acoustic modes around 0.5 THz and the lowest-energy optic branches around 2.0 THz, which both have the most negative $\gamma_{k,\lambda}$ and span the entire Brillouin zone. This was also highlighted in the DoS in Fig. 11(a), where we coloured the DoS according to the mean values of $\gamma_{k,\lambda}$ for each frequency bin. Then, we calculated the rigid unit modes (RUMs) of Zn(CN)$_2$ using the CRUSH code\cite{13} as the set of eigenvectors of the dynamical matrix whose

**FIG. 8.** Vibrational DoS calculated using MD at 300 K and at 0.026 GPa, 0.20 GPa, 0.46 GPa and 0.65 GPa. (a) shows the full DoS for the CN-disordered model; (b) shows the full DoS for the CN-ordered model; (c) shows the DoS for only C–N rigid rod rotations. The peaks at around 14 THz, corresponding to pure Zn–C(N) bond flexing, increase in frequency on compression.

**FIG. 9.** Vibrational DoS calculated using MD at 0.0 GPa and temperatures 162 K, 297 K, 473 K and 603 K. (a) shows the full DoS; (b) shows the DoS for only C–N rigid rod rotations. The peaks in the full DoS at around 14 THz, corresponding to pure Zn–C(N) bond flexing, broaden and decrease in frequency on heating.

pure Zn–C/N bond flexing — it completely disappears in the angular DoS. Unlike the acoustic peak, this peak is softened on heating due to the thermal expansion of the Zn–C/N bond, and flattens with elevated temperature due to the finite life time of the corresponding phonon.
FIG. 10. (a) Dispersion curves coloured according to the corresponding value of the mode Grüneisen parameter \( \gamma_{k,\lambda} \). Red is a negative value of \( \gamma_{k,\lambda} \) (down to a minimum of \(-18\)); blue is a positive value of \( \gamma_{k,\lambda} \) (up to a maximum of 0); white is a \( \gamma_{k,\lambda} \) value of zero. (b) Dispersion curves coloured according to projection of the eigenvectors onto RUMs. Colour strength corresponds to the degree to which a mode is a RUM; white corresponds to zero RUM character in a given mode. The colour itself corresponds to the nature of the RUM: Red is purely rotational RUM motion; blue is purely translational RUM motion. The corresponding eigenvectors of (c) the translational RUMs (around 0.5 THz), (d) the first rotational RUMs (around 2.0 THz) and (e) the second rotational RUMs (around 9.0 THz), viewed down the [1,1,0] direction with the undistorted structure in grey shown behind. The translational RUMs correspond to lateral translations of the tetrahedra leading to rotations of pairs of tetrahedra. The rotational RUMs correspond to neighbouring tetrahedra rotating in the same or opposite direction.

FIG. 11. Full DoS calculated using lattice-dynamics. (a) Each bin is coloured according to its average Grüneisen parameter: red bins have an average \( \gamma_{k,\lambda} \) value of \(-10\); white bins have a positive average \( \gamma_{k,\lambda} \). (b) Each bin is coloured according to its average RUM component. Red bins are pure RUMs; blue bins have zero RUM character.

FIG. 12. Profiles of mode Grüneisen parameters of Zn(CN)_2 calculated at three different pressures from lattice dynamics: black curve is 0.0 GPa; blue curve is 0.2 GPa; red curve is 0.4 GPa. The Grüneisen parameters of the modes around 0.5 THz, 2.0 THz and 9.0 THz become more negative as elevated pressures, resulting in a more negative overall Grüneisen parameter as pressure is increased.

eigenvalues are zero. We took the dot product between the eigenvectors of the RUMs and the eigenvectors from the lattice-dynamic calculation, and then coloured the dispersion curves in Fig. 10(b) by the extent to which each mode eigenvector can be described in terms of correlated whole-body translations (in blue) and rotations (in red) of [Zn(C/N)_4] tetrahedra. The DoS in Fig. 11(b)
metrical Anharmonicity ($K^\text{a}$) with frequencies less than 1 THz ($\sim$ 0.5 THz) shows the largest value compared to others. The modes around 15 THz corresponding to Zn–C(N) bond flexing show negative anharmonicity. The anharmonicity of all these modes is enhanced by compression.

was also coloured accordingly. One can see that all the modes with negative $\gamma_{k,\lambda}$ that contribute to the NTE of the material are RUMs.

As shown by Fig. 10(c), the acoustic modes around 0.5 THz, like those at M and X, are characterized by translational motions of the rigid tetrahedral units, partly involving angular rotations of the C–N rod. The optic modes around 2.0 and 9.0 THz can be seen as neighbouring tetrahedral rotating against each other, as respectively shown by Fig. 10(d) and (e). The RUM nature of these modes guarantees their low frequencies and large negative $\gamma_{k,\lambda}$. The relatively high frequencies of the optic RUMs is due to the breaking of the Zn–C/N–N/C alignment, and the magnitudes of their negative $\gamma_{k,\lambda}$ suffer accordingly. Study of the eigenvectors also directly revealed that the non-RUM modes around 10 and 15 THz correspond to the pure angular vibration of C/N–Zn–N/C within the tetrahedra and the pure bond flexing of Zn–C/N, respectively.

We also found that, besides the negative $\tilde{\gamma}$, Zn(CN)$_2$ also has negative $\partial \tilde{\gamma} / \partial P$ and $\partial^2 \tilde{\gamma} / \partial P^2$. Fig. 12 shows the profiles of $\gamma_{k,\lambda}$ up to 0.4 GPa calculated in HLD. It is clear that $\tilde{\gamma}$ becomes more negative on compression due the contributions from the RUMs around 0.5 THz, 2.0 THz and 9.0 THz, with the translational RUMs having the most negative $\gamma_{k,\lambda}$ (refer to Fig. 11(a)) contributing the most. The negativity of the pressure derivatives of $\tilde{\gamma}$ would result in negative $B'_0$ at non-zero temperatures.

C. Anharmonicity and quantum effects

We found that the acoustic peak in the DoS (see Fig. 11) with frequencies less than 1 THz ($\sim$ 50 K) to be the most negative. One can see that all the modes with negative $\gamma_{k,\lambda}$ that contribute to the NTE of the material are RUMs.

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C. Anharmonicity and quantum effects

We found that the acoustic peak in the DoS (see Fig. 11) with frequencies less than 1 THz ($\sim$ 50 K) ac-
FIG. 15. Temperature dependence of the normalized anharmonicity at 0.0 GPa. At 25, 280 and 560 K, calculations were conducted using the cumulative distributions of couples of DoS at 25/45 K, 280/300 K and 560/590 K, respectively. The general trend is that the mode frequencies change less rapidly with temperature on heating.

inclined in the formalism of Eq. 1 to calculate $\alpha_V$. Together with the direct MD and QHLD results, the temperature dependence of the NTE from this method (‘MD+QE’) is shown in Fig. 13. At low temperatures, the curve acts like the QHLD result due to quantum quenching. At high temperatures, the curve becomes less negative like MD due to anharmonicity, making the curve in a better agreement with the experiment.

From the same analysis, we can also obtain the temperature and pressure dependence of anharmonicity. The normalized anharmonicity, measuring the change of mode frequency with temperature at constant pressure is defined as

$$\frac{1}{\omega_s} \frac{\partial \omega_s}{\partial T} \bigg|_p - \gamma_s \alpha_V,$$

where, on the right-hand side, the first term is the intrinsic anharmonicity and the second term is the contribution from the contraction of lattice (implicit). By using the DoS from the MD at 180 and 240 K, the normalized anharmonicity of each mode was calculated, as shown in Fig. 14(a). The results of the important modes around 0.5, 2.0 and 9.0 THz, as indicated by the vertical lines, agree quite well with the experiment (using DoS at the 180 and 240 K).

The implicit anharmonicity was calculated using the mode Grüneneise parameters and the ‘MD+QE’ value of $\alpha_V$ at 180 K. The intrinsic anharmonicity was then obtained from Eq. 4.

The normalized anharmonicity at different pressures is shown in Fig. 14(b). As mentioned in the former sections, the modes around 0.5, 2.0 and 9.0 THz, corresponding to the translational and rotational RUMs, respectively, are stiffened on heating with positive normalized anharmonicity. Among these, the translational RUMs (around 0.5 THz) show the largest normalized anharmonicity of more than $4 \times 10^{-4}$ K$^{-1}$ at 0.0 GPa. The modes around 15 THz corresponding to pure bond flexing of Zn–C(N) are softened on heating with negative anharmonicity. The normalized anharmonicity of all these peaks is strengthened on compression.

We further calculated the normalized anharmonicity at low (25 K), medium (280 K) and high (560 K) temperatures to see its temperature dependence, as shown in Fig. 15. At each temperature, two DoS with temperature difference less than 30 K are used. The low-temperature (25 K) value of the anharmonicity of the modes around 0.5 THz is $1.0 \times 10^{-3}$ K$^{-1}$, in good agreement with the experimental value of $1.1 \times 10^{-3}$ K$^{-1}$ in Ref. 41. The figure also shows that the mode frequency would change less rapidly on heating at high temperatures. The same trend is seen for the 0.5 THz peak in the experiment in Ref. 41.

VIII. STRUCTURAL STUDY

A. The local picture on compression and heating

In real space, variations of geometrical features of the material, such as the Zn–C/N bond length, the N/C–Zn–C/N angle within the tetrahedral unit, and the Zn–C/N–N/C angle are important for us to build a local picture...
of the system subject to both compression and heating. Distributions of these quantities, as shown in Fig. 10, were obtained from the atomic trajectory data of the MD. The large spread of the distributions suggests large vibrations of these quantities at high temperature. The slightly expansion-biased broadening of the bond-length distribution on heating indicates an enhanced thermal expansion in the bond. We found that the deviation of the average N/C–Zn–C/N angle in the tetrahedral unit from its equilibrium of 109.47° is trivially small (∼0.2°) even at very high temperature (∼600 K) and pressure (∼0.6 GPa).

The average bond length and the average angle distortion in Zn(CN)₂ as functions of both temperature and pressure are shown in Fig. 17 and 18, respectively. Compression progressively increases the Zn–C/N–N/C angle with elevated temperature. At very low and zero temperature, the trend is that the angle will be hardly changed by pressure. This is exactly what expected in the previous section where we suggested that, at zero temperature, the volume change of the material due to the pressure arises solely from the compression of the Zn–C/N bonds, resulting in positive B₁₀. It is when the Zn–C/N–N/C angle starts to increase under compression and contribute to the volume change that the material shows negative B₁₀, i.e. the pressure-induced softening. The inset of Fig. 17 shows the increase of bond length with temperature. The superlinear behaviour indicates the softening of the bond at higher temperature due to the thermal expansion. In Fig. 18, the inset shows the increase of the Zn–C/N–N/C angle with elevated temperature. The sublinear behaviour suggests that the angle will become more rigid, which should result in a less negative B₁₀ at high temperatures. Both the superlinearity and the sublinearity in the plots rely on the capture of the anharmonicity of the material.

To see the real-space picture of RUMs in Zn(CN)₂, we quantified the proportion of thermal-excited rigid-unit rotations at different pressures. Using our GASP code, we compared ten snapshots of an MD simulation, where each snapshot is separated by 2 ps, with the ideal structure. This is repeated for various pressures and temperatures. GASP, using geometric algebra, can partition the atomic displacements for every comparison made into the mean squared rigid-tetrahedron rotations, translational displacements and unit deformations. We then computed the average proportion of rigid-unit rotations at each temperature. To exclude those rigid-unit rotations due to pure topology reasons, i.e. rotations that accidentally maintain the shape of the tetrahedral unit under thermal excitation but are not because of the features of motion, we set up a benchmark calculation using an ideal cristobalite structure without any interactions other than bonds to hold Zn–C/N and C–N. The reason to use the single-framework lattice is to avoid the problem of two interpenetrating frameworks crushing into each other in the MD due to the lack of long-distance interactions. Fig. 19 shows the results at different pressures and temperatures with coloured areas. The proportion of the rigid-unit rotation of a real silica system is given in the plot as a comparison. The figure suggests that compression will enhance the rigid-unit rotations. At ambient temperature ∼300 K, for example, the average proportions are 63% at 0.0 GPa, 64% at 0.2 GPa, 65% at 0.4 GPa and 66% at 0.6 GPa, compared to 35% of the benchmark and 90% of the silica system. Another important point is that, at certain pressure, the proportion...
FIG. 19. Plot showing the proportion of rotational RUMs present in Zn(CN)$_2$ as a function of temperature, calculated using GASP and data from MD simulations. Data is presented for four pressures plus two additional benchmark systems. The green region is the rotational component present in the ideal cristobalite structure; the yellow region is the additional rotational component present in Zn(CN)$_2$ at 0.0 GPa; the orange region is the additional component present at 0.2 GPa, the pink region is the additional component at 0.4 GPa; the red region is the additional component at 0.6 GPa and the dark red region is the additional component present in amorphous silica. The results clearly show the enhancement of the rotational RUMs in Zn(CN)$_2$ under pressure, as well as the reduction of the rotational RUMs on heating.

of rigid-unit rotation will decrease with elevated temperature. This trend corresponds to the peaks around 0.5, 2.0 and 9.0 THz in DoS stiffened on heating, as shown in Fig. [3]. The Grüneisen parameters of these modes will consequently become less negative, and so does the coefficient of thermal expansion.

B. The nearest-neighbour Zn$_i$–Zn$_j$ distance

There is a puzzling observation from the X-ray pair distribution function measurements of Zn(CN)$_2$ that instantaneous Zn$_i$–Zn$_j$ distances contract less rapidly on heating than does the cell length, while crystallographically the two should be linked. The explanation relies on the nature of the acoustic modes as translational RUMs.

The acoustic modes around 0.5 THz are translational RUMs which count for half of the NTE of the material as mentioned in Sec. VII. Unlike rotational RUMs around 2.0 and 9.0 THz that will reduce the nearest Zn$_i$–Zn$_j$ distance, the translational RUMs correspond to collective translations of the neighbouring rigid units that moves zinc atoms off site and retains the distance of the nearest-neighbour zines, as seen in Fig. [10(c)]. This kind of vibration involves the rotation of C–N rod around its centre of mass, consistent with the previous finding in the DoS that part of the acoustic modes are from the C–N rod rotations.

We calculated the average distance of the nearest Zn$_i$–Zn$_j$ using the the trajectory data from the MD simulations. Fig. [20] shows the temperature dependence of the average distance under different pressures. At zero pressure, the ratio between the linear coefficient of thermal expansion (CTE) of the averaged Zn$_i$–Zn$_j$ distance, $\alpha_{ZnZn}$, and the overall linear CTE of the material, $\alpha$, is 0.67, with $\alpha_{ZnZn} = -8.5\text{MK}^{-1}$ and $\alpha = -12.62\text{MK}^{-1}$. This result agrees well with the experimental ratio of 0.71, with $\alpha_{ZnZn} = -12.42(12)\text{MK}^{-1}$ and $\alpha = -17.40(18)\text{MK}^{-1}$.

IX. DISCUSSIONS AND CONCLUSIONS

This study shows that almost all the modes responsible for the NTE of Zn(CN)$_2$ are RUMs. We managed to categorize these modes in terms of their vibrational motions. The TA modes around 0.5 THz spanning to the lowest energy correspond to collective motions of Zn–CN–Zn as a rigid body, which can keep the distance of the nearest-neighbour zins. The low energy hence the most negative Grüneisen parameters of these modes are due to this kind of collective motion without involving relatively high-energetic angle bending in the Zn–CN–Zn linkage. These modes contribute half of the NTE of the material. The optic modes around 2.0 THz and 9.0 THz correspond to rotations of the neighbouring tetrahedral units against each other involving angle bending in the Zn–CN–Zn linkage, resulting in higher mode energy and...
less negative Grüneisen parameters.

Although the increase of pressure or temperature would both result in volume contraction in Zn(CN)₂, the pressure and temperature dependence of the NTE in Zn(CN)₂ are totally different. Increasing temperature stiffens the low-frequency peaks and softens the high-frequency peaks in the DoS (Fig. 9 and Fig. 14) accompanied by the reduction of NTE, while compression would soften the low-frequency peaks and stiffen the high-frequency peaks (Fig. 8) resulting in NTE enhancement. Raising temperature slows the mode softening caused by compression, and postpones the phase transition. The enormous decrease of the bulk modulus on heating contrasts the small change of the bulk modulus on compression, which disobeys Birch’s law of corresponding states.

The pressure and temperature dependence of the Zn–C/N bond and the N/C–Zn–C/N angle are also intriguing. The large vibrational amplitude of the N/C–Zn–C/N angle extends the Zn–C/N bond, which can be seen (under constant pressure) in Fig. 17. The superlinearity suggests an enhancement of the thermal expansion in the bond with more distorted angle at higher temperature. The sublinearity of the average angle distortion corresponds to the stiffened RUMs involving the rotational vibrations of the C–N rods.

The ability to carry out MD for Zn(CN)₂ using the potential model is vital in this study. It allows us to capture the anharmonicity to reproduce the exotic properties of the material, and to study their pressure and temperature dependence. The origins of various properties are revealed by linking features in both energy and real space.

One example is the pressure-enhanced NTE, which has been well reproduced in Fig. 8(a). This behaviour is a natural result followed by softening of the bulk modulus on heating (Eq. 2), and is linked to the feature in energy space that the modes around 0.5, 2.0 and 9.0 THz are softened under compression (as in Fig. 8), as well as the rising proportion of the rigid-unit rotations in real space on compression (as in Fig. 19).

Another example is the reduction of NTE with elevated temperature. stiffening of the modes around 0.5, 2.0 and 9.0 THz on heating (as shown in the DoS in Fig. 9) makes their Grüneisen parameters less negative, hence the reduction of NTE in the material. In real space, Fig. 19 clearly shows the trend of reduction in the proportion of rigid-unit rotations on heating.

The third example is the temperature dependence of the bulk modulus and its first derivative B"₀. The large decrease of the bulk modulus on heating is due to the involvement of the Zn–C/N–N/C angular vibrations at non-zero temperature and the softening of the Zn–C/N bond, which can be seen both in the DoS (Fig. 9(a)) and in the real-space picture of Fig. 16, 17 and 18. As shown in Fig. 18, the average Zn–C/N–N/C angle starts to possess an almost linear pressure dependence at medium temperature and contributes to the pressure-induced volume change of the material. According to a simple geometrical relation, the relative decrease in cell is roughly proportional to the Zn–C/N–N/C angle squared, hence to the pressure squared, resulting in negative B"₀. However, this contribution from the pressure-induced change in the Zn–C/N–N/C angle to the volume contraction will be hindered at high temperature due to the anharmonic stiffening of the corresponding modes (Fig. 9(a) and Fig. 14). On the other hand, the contribution from Zn–C/N bond compression will increase due to the thermal softening of the bond. The combined effect of these two trends is expected to result in a less negative B"₀ at high temperatures.

ACKNOWLEDGMENTS

We gratefully acknowledge financial support from the Cambridge International Scholarship Scheme (CISS) of the Cambridge Overseas Trust and Fitzwilliam College of Cambridge University (HF), NERC and CrystalMaker Software Ltd. (LHNR). The MD simulations were performed using the CamGrid high-throughput environment of the University of Cambridge. The interatomic potential was developed through our membership of the UK HPC Materials Chemistry Consortium, funding by EPSRC (EP/F067496), using the HECToR national high-performance computing service provided by UoE HPCx Ltd at the University of Edinburgh, Cray Inc and NAG Ltd, and funded by the Office of Science and Technology through EPSRC’s High End Computing programme.

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