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Phonon transport and thermal conductivity of diamond superlattice nanowires: a comparative study with SiGe superlattice nanowires

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Due to the coupling of a superlattice's longitudinal periodicity to a nanowire's radial confinement, the phonon transport properties of superlattice nanowires (SLNWs) are expected to be radically different from those of pristine nanowires. In this work, we present the comparative investigation of phonon transport and thermal conductivity between diamond SLNWs and SiGe SLNWs by using molecular dynamics simulations. In the case of period length ~ 25 Å, the thermal conductivities of diamond SLNWs and SiGe SLNWs both increase linearly with increasing the period number, which implies the wave-like coherent phonons dominate the heat transport of SLNWs. In the case of period length ~ 103 Å, the thermal conductivity of SiGe SLNWs is length-independent with increasing the period number, indicating that the particle-like incoherent phonons in SiGe SLNWs control the heat transport, because the phonon–phonon scattering causes phonons to not retain their phases and the coherence is destroyed before the reflection at interfaces. However in diamond SLNWs the coherent phonons still dominate heat conduction and the thermal conductivity is length-dependent, because the mean free path of phonon–phonon scattering in diamond SLNWs is much longer. The spatial distribution of phonon localized modes further supports these opinions. These results are helpful not only to understand the coherent and incoherent phonon transport, but also to modulate the thermal conductivity of SLNWs.

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1. Introductions

Diamond is the stiffest material in nature, and has extremely high thermal conductivity which can exceed 2000 W m⁻¹ K⁻¹ at room temperature. The thermal transport properties of pristine diamond,1 diamond nanothread,2 isotope-doped diamond3 and diamond composites4 have been widely investigated. It is well known that diamond has hexagonal and cubic structures. Tanigaki et al.5 reported that a nanopolycrystalline diamond was synthesized from graphite by a direct-conversion method. They found that nanopolycrystalline diamond was a superlattice-like twinned structure composed of hexagonal and cubic diamonds. Meanwhile, by ab initio-calculations they demonstrated that the nanopolycrystalline diamond was stiffer than the natural and synthesized monocrystal diamond. Plenitudinous studies focused on the structures and mechanical properties of nanopolycrystalline diamond,6-9 however the research of phonon transport properties of this novel structure is still scarce so far.

Phonon transport and thermal conductivity of nanomaterials have attracted considerable research effort due to

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their unique properties and potential applications, 10-20 such as thermoelectric generators and thermal management devices. Generally, there are two contrasting pictures to describe phonon transport in nanostructures, the one is particle-like incoherent phonons, and the other is wave-like coherent phonons.21 Recently, the influence of coherent and incoherent scattering mechanisms on thermal conductivity of periodic structures, such as superlattices and nanomeshes, is one of the hottest topics in the field of nanoscale heat transport.22-31 In periodic nanostructures, if the mean free path of phonons is longer than the period length of structures, coherent phonon transport occurs due to the constructive interference of incident and reflected phonon waves at the interfaces. 32,33 At room temperature, phonons in diamond have long mean free path, because C-C bond is very strong and the phonon-phonon Umklapp scattering in diamond is weak.34 Therefore, superlattice-like structures of diamond, such as nanopolycrystalline diamond, are beneficial for investigating coherent phonon transport mechanism.

In this work, we demonstrate the strongly length-dependent thermal conductivity of diamond SLNWs constructed from hexagonal and cubic diamond by non-equilibrium molecular dynamics (NEMD) simulation at room temperature, which implies coherent phonon transport. There are many factors that affect the phonon coherence in superlattice structures, and the interface is one of the most important. In order to investigate the

coherent and incoherent phonon transport at the interface, the thermal conductivities of diamond SLNWs and SiGe SLNWs are comparatively studied. The calculation results show that the thermal conductivities of diamond SLNWs and SiGe SLNWs have the same linearly increasing trend with the number of period, when the period length is $\sim\!\!25$ Å. The localization effect of phonons is not observed at the interfaces of both SLNWs. When the period length is $\sim\!\!103$ Å, the thermal conductivity of diamond SLNWs still increases with the number of period. No localized effect of phonons is observed at the interface. However, the thermal conductivity of SiGe SLNWs converges to a constant, and obvious phonon localization effect is observed at the interface. This indicates that phonon transport is still coherent in diamond SLNWs, while it is incoherent in SiGe SLNWs.

Models and methods

Fig. 1(a) shows the structure of diamond SLNW, A' and B' are the structure of hexagonal diamond, the A, B and C are the structure of cubic diamond. The diamond SLNW is composed of cubic and hexagonal diamond alternately stacked in the *z*-axis direction. The lattice constant is 2.058 Å. Fig. 1(b) shows the cross-section of diamond SLNW, and Fig. 1(c) shows the model of diamond SLNW. In Fig. 1(b) and (c), the red part represents cubic diamond, and the blue part represents hexagonal diamond. The ratio of hexagonal diamond to cubic diamond is $1:1.\ L_s$ is the period length of SLNW in Fig. 1(c).

In this work, the non-equilibrium molecular dynamics (NEMD) simulation method is used to calculated the thermal conductivity of SLNWs. We adopt the Tersoff potential³⁵ for both carbon, silicon, and germanium atoms. All simulations are performed using the LAMMPS³⁶ code with a time step of 0.5 fs. As shown in Fig. 2, we divide the sample into three parts: fixed wall at each end of the sample, thermostat area adjacent to the fixed wall, and the free part in the center. An fixed wall is used to make the atoms in the thermostat areas steady, and equalize the force applied on them. Free boundary conditions are used in all directions.

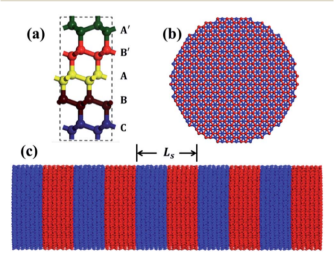


Fig. 1 (a) The structure of diamond SLNW. (b) The cross section of diamond SLNW. (c) The vertical axis plane of diamond SLNW.

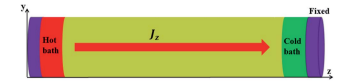


Fig. 2 Simulation model of thermal conductivity of SLNWs.

In our simulations, using a Nosé-Hoover thermostat^{37,38} we firstly equilibrate this system at 300 K for 5 ns with the walls moving freely in the longitudinal direction, corresponding to zero pressure. After the NPT relaxation, the optimized structure is obtained, we fix the atoms at the ends of the sample and run the system in the constant volume and constant temperature ensemble (NVT) for 2 ns. After the NVT, the system continues to be simulated in the constant volume and constant energy ensemble (NVE) for 2 ns. After these running, to obtain steady heat flux, the temperature of hot and cold Nosé-Hoover thermostat is set as 320 K and 280 K respectively. Due to the existence of temperature difference, heat flows from the hot bath to the cold bath across the sample. The steady state is reached after this NEMD is performed for 5 ns. The simulation of final 2 ns is to get the heat flux and calculate the thermal conductivity. The temperature gradient (dT/dz) is obtained by fitting the linear portion of temperature profile, and the heat flux (I_z) is calculated using:

$$J_{z} = \frac{1}{A} \sum_{i \in \text{hot bath}} \frac{\mathrm{d}}{\mathrm{d}t} E_{i}(t) \tag{1}$$

where E is the energy, t is the simulation time (2 ns), and A is the cross-section area. To calculate the cross-section area, we approximate the cross-section of SLNWs to a circle. The thermal conductivity κ is then calculated by using Fourier's law of heat conduction:

$$\kappa = -J_z/(\mathrm{d}T/\mathrm{d}z) \tag{2}$$

Results and discussions

The thermal conductivities of the diamond SLNWs as a function of the number of periods are shown in Fig. 3(a). In the case of short period length such as $L_{\rm s}=24.7$ Å. The thermal conductivity is almost linearly proportional to the number of periods. This ballistic transport across the whole thickness implies coherent heat conduction in the diamond SLNWs. The nonlinear dependence of thermal conductivity on the number of periods in the case of long period length $L_{\rm s}=102.9$ Å suggests the increasing influence of incoherent effects. However, the thermal conductivity is still obviously increasing with the increase of period number, therefore coherent phonons still dominate the heat conduction in diamond SLNWs.

For comparison, we also calculate the thermal conductivities of the SiGe SLNWs with the same period length as a function of their number of periods, which is shown in Fig. 3(b). It is evident that the thermal conductivity of SiGe SLNWs with $L_{\rm s}$ of \sim 25 Å

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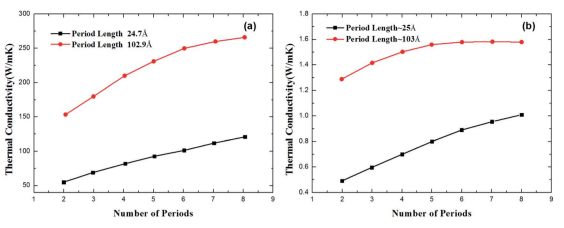


Fig. 3 Thermal conductivities of diamond SLNWs (a) and SiGe SLNWs (b) versus the number of periods

increases almost linearly with the number of periods, implying coherent phonon transport characteristics. However, in the case of long period length ~ 103 Å, the situation is quite different. When the number of periods is larger than 5, the thermal conductivity of SiGe SLNWs is length-independent, indicating that the incoherent phonons dominate the thermal transport.

In order to further reveal the difference of phonon transport mechanisms between diamond SLNWs and SiGe SLNWs, we investigate the localization effect of the phonon modes at the interface in these two types of SLNWs with the $L_{\rm s}$ of ~ 103 Å. The position of localized phonons can be described clearly by the spatial distribution of localized modes $\Gamma = \{\lambda : p_{\lambda} < p_{\rm c}\}$. The spatial distribution of localized modes is expressed by

$$\varnothing_{\Gamma}(i) = \frac{\sum_{\lambda \in \Gamma} \sum_{\alpha} \varepsilon_{i\alpha,\lambda}^* \varepsilon_{i\alpha,\lambda}}{\sum_{i} \sum_{\lambda \in \Gamma} \sum_{\alpha} \varepsilon_{j\alpha,\lambda}^* \varepsilon_{j\alpha,\lambda}}$$
(3)

where, p_c is a criteria for localization. Zhang *et al.* considered the phonon mode was fully localized if the participation ratio of this phonon vibrational mode was lower than 0.2,³⁹ so we choose the criteria $p_c = 0.2$. p_{λ} is the phonon vibration mode participation ratio, which is given by

$$P_{\lambda}^{-1} = N \sum_{j} \left(\sum_{\alpha} \varepsilon_{j\alpha,\lambda}^* \varepsilon_{j\alpha,\lambda} \right)^2$$
 (4)

here, $\varepsilon_{j\alpha,\lambda}$ denotes the α -th vibrational eigenvector component of the normal eigen-mode λ for the jth atom, α denotes the Cartesian direction, j sums over all the atoms. The value of $\mathcal{O}_{\Gamma}(i)$ denotes the degree of localization at position (X, Y, Z) of atom i.⁴⁰

The normal-mode eigen frequencies and their corresponding eigenvector components are gotten *via* solving the lattice dynamical equation expressed by

$$\omega_{\lambda}^{2} \varepsilon_{i\alpha,\lambda} = \sum_{i\beta} \Phi_{i\alpha,j\beta} \varepsilon_{j\beta,\lambda} \tag{5}$$

where the elements of the force constant matrix Φ are given as:

$$\Phi_{i\alpha,j\beta} = \frac{1}{\sqrt{m_i m_j}} \frac{\partial^2 V}{\partial u_{i\alpha} \partial u_{j\beta}} \tag{6}$$

here $u_{i\alpha}$ is the displacement of atom i in the α Cartesian direction, m_i is the mass of atom i, and V is the total potential energy of the system.

Fig. 4(a) and (b) demonstrate the spatial distribution of localized phonon modes in diamond SLNWs within $\Gamma = \{\lambda : p_{\lambda} < 1\}$ 0.2} at the interface and non-interface atom layers, respectively. It is obvious that no matter interface or non-interface, the localized phonon modes locate in the surface region of diamond SLNWs. Chen et al. found that because the dangling bond atoms on the surface break the perfect lattice periodicity, phonon modes are localized on the surface of nanowires.41 Except for atoms in the surface region, the inner atoms have ultralow value of \emptyset_{Γ} , and there are no significant differences for the spatial distribution of localized modes between the interface and non-interface atom layers. This implies that the phonons are not localized at the interface, and the coherent phonons transport through the interface in the diamond SLNWs with $L_{\rm s}$ of \sim 103 Å. The analysis agrees well with the theoretical and experimental results,32,42,43 in which coherent phonons maintain their phase when they cross multiple interfaces in a superlattice, and the superlattice can be considered as a homogeneous material with its own primitive cell and phonon spectra.

The spatial distribution of localized phonon modes for SiGe SLNWs within $\Gamma=\{\lambda:p_{\lambda}<0.2\}$ at the interface and non-interface atom layer is shown in Fig. 5(a) and (b) respectively. The spatial distribution of localized modes at the interface and non-interface layer is quite different. Despite the surface region, phonons propagate through the non-interface layer, but they are strongly localized at the interface layer of SiGe SLNWs. From Fig. 4(b) and 5(b), we consider that there are two competing mechanisms controlling the heat transport in SLNWs, which are particle-like interface scattering and wave-like interference effect. The thermal conductivity of diamond is ultra-high. According to the kinetic theory $\kappa=\frac{1}{3}Cvl$ Che *et al.* derived the phonon mean free path l=494 nm at room temperature. Due to the large mean free path, phonon–phonon Umklapp scattering is scarce between the reflection at atomically smooth interfaces, a majority of phonons in diamond SLNWs can maintain their phases, and the wave-like interference effect

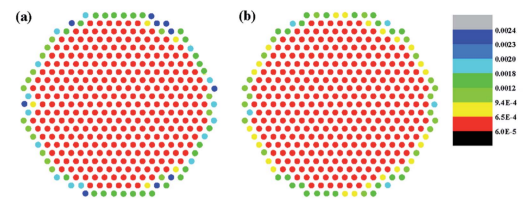


Fig. 4 Spatial distribution of localized phonon modes in diamond SLNWs at the interface atom layer (a), and at the non-interface atom layer (b). Positions of the circles denote the atom locations on the XY plane, and spatial distribution of localized phonon modes is depicted according to the colour.

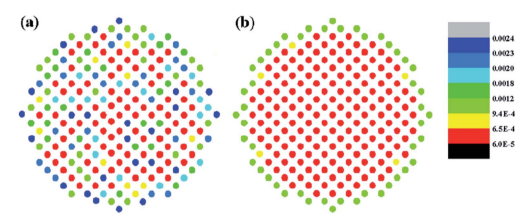


Fig. 5 Spatial distribution of localized phonon modes in SiGe SLNWs at the interface atom layer (a), and at the non-interface atom layer (b).

dominates heat conduction in diamond SLNWs. However, the thermal conductivity of silicon and germanium is much lower than diamond and accordingly the phonon mean free path of silicon is much shorter. The frequent phonon-phonon scattering make phonons not retain their phases and the coherence is destroyed before the reflection at interfaces, therefore phonons in SiGe SLNWs are particle-like, and the interface scattering mechanism is much stronger than the coherent transport mechanism. It is worth noting that in the case of very short period length, phonon-phonon Umklapp scattering is scarce between the reflection at interfaces, therefore coherent transport mechanism is dominant in SiGe SLNWs, and the thermal conductivity is evidently length-dependent, shown by the black line in Fig. 3(b).

4. Conclusions

In summary, we calculate the thermal conductivities of diamond SLNWs and SiGe SLNWs by performing the non-equilibrium molecular dynamics simulations, and investigate the phonon transport mechanisms in SLNWs. We find that there are two competing mechanisms controlling the heat transport in SLNWs, which are particle-like interface scattering

and wave-like interference effect. Which mechanism is dominant depends on the relationship between phonon-phonon scattering mean free path and period length of SLNWs. In the case of short period length (\sim 25 Å), the phonon-phonon scattering mean free paths of both SLNWs are longer than the period length, the phonons can retain their phases and wave nature, therefore the heat transport is coherent, and the thermal conductivities are linearly length-dependent. In the case of long period length (\sim 103 Å), a majority of phonons in SiGe SLNWs can not maintain their phases because of frequent phonon-phonon scattering, and the coherence is destroyed. Phonons are strongly localized at the interface layer of SiGe SLNWs, indicating that the particle-like interface scattering mechanism is dominant, and the interfaces significantly hinder the heat conduction, therefore the thermal conductivities are length-independent. However, due to the ultra-long phononphonon scattering mean free path of diamond SLNWs, in this case (period length \sim 103 Å), the wave-like interference effect still dominates the heat conduction, and the phonon transport is quasi-ballistic. Our findings are helpful not only for understanding the coherent and incoherent phonon transport mechanism, but also for manipulating the thermal conductivity of SLNWs.

Conflicts of interest

The authors declare no conflict of interest.

Acknowledgements

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