Heat conduction of single-walled carbon nanotube isotope superlattice structures: A molecular dynamics study

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Heat conduction of single-walled carbon nanotubes (SWNTs) isotope superlattice is investigated by means of classical molecular dynamics simulations. Superlattice structures were formed by alternately connecting SWNTs with different masses. On varying the superlattice period, the critical value with minimum effective thermal conductivity was identified, where dominant physics switches from zone-folding effect to thermal boundary resistance of lattice interface. The crossover mechanism is explained with the energy density spectra where zone-folding effects can be clearly observed. The results suggest that the critical superlattice period thickness depends on the mean free path distribution of diffusive-ballistic phonons. The reduction of the thermal conductivity with superlattice structures beats that of the one-dimensional alloy structure, though the minimum thermal conductivity is still slightly higher than the value obtained by two-dimensional random mixing of isotopes.

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I. INTRODUCTION

The ever-expanding expectations for single-walled carbon nanotubes (SWNTs) (Ref. 1) include applications for various electrical and thermal devices due to their remarkable electrical and thermal properties.2,3 On considering the actual applications, one of the essential tasks is to characterize the thermal properties certainl for thermal devices and also for electrical devices, since they limit the affordable amount of electrical current through the system. In actual applications, SWNTs may have impurities, defects, and junctions, which alter the heat conduction. In general, these nanoscale impurities, having scales comparable to the phonon mean free path $l$ of the system, are expected to strongly influence the thermal properties of bulk materials. For SWNTs, due to the quasi-one-dimensional structure and the expected long $l$, the impact and its scale dependence of the impurity effects may differ considerably from those of the three-dimensional materials. In our previous molecular dynamics study, isotope effects on the thermal conduction were investigated by randomly mixing $^{13}$C isotopes in a finite length $^{12}$C-SWNT with chirality of (5, 5).4 The results showed that the effective thermal conductivity decreases monotonically as the number ratio of $^{13}$C atoms increases. Interestingly, the isotope effect was found to be considerably weaker than that in diamonds. This led us to speculate that the phonon impurity scattering may be restricted in SWNTs due to the one-dimensional confinement of phonons.

In the current work, we proceed with the characterization of heat conduction of SWNTs subjected to nanoscale intrinsic thermal resistances. Here, in order to reduce the complication due to the disorderliness of impurities in the previous work,4 we investigate the heat conduction of SWNTs altered by systematically placed thermal resistances on well-defined scales. To this end, the superlattice structures were constructed by periodically connecting SWNTs with different masses. There have been a considerable number of studies devoted to characterize thermal properties of superlattice structures in three-dimensional materials. Being motivated by thermoelectric device applications, the main purpose of these studies has been to utilize the superlattice structures to reduce the effective thermal conductivity $\lambda$ in the direction normal to the interfaces. The challenge has been to beat $\lambda$ of alloys (alloy limit), which classically, among binary crystal materials, has been considered to possess the minimum thermal conductivity. Experimental and theoretical studies of the cross-plane thermal conductivity and its dependence on the period thickness have been reported for thin film superlattices5–12 and recently for nanowire superlattices.13–15 The most common experimental observation has been the reduction of thermal conductivity on shortening the period thickness $\Delta z$ (Ref. 5) and hence extensive theoretical studies have been directed to reproduce such trend.6 On the other hand, experiments of short-period superlattices revealed the existence of the minimum thermal conductivity for a period thickness of about 5 nm.7,8 This was explained by a study using molecular dynamics models, where the minimum thermal conductivity was observed for a period thickness of several monolayers.9

The appearance of the minimum thermal conductivity can be understood in terms of two competing mechanisms that yield opposite dependence of $\lambda$ on $\Delta z$.10 One is the zone-folding effect due to new periodicity imposed by the superlattice structure. The reduction of the group velocity and enhancement of umklapp scattering at the imposed zone-boundaries (mini-Brillouin zone boundaries), which appear as the gaps in the dispersion relations (mini band gaps), attenuate the thermal conductivity.11,12 The effect becomes stronger as $\Delta z$ increases, since the number of folds is proportional to the period thickness. The other mechanism is the thermal boundary resistance (TBR) at the superlattice interfaces due to phonon reflection and scattering. The TBR effect increases with the number of interfaces per length and hence decreases with $\Delta z$. The reduction of $\lambda$ attributed to TBR typically saturates on reducing $\Delta z$ below the characteristic length of the ballistic phonon transport due to the phonon tunneling effects, i.e., the transboundary ballistic transport of phonons through TBRs. Simkin and Mahan,10 in
their quantum mechanical model, implemented the crossover of the two competing physics by using the imaginary wave vector. The results led them to suggest that the critical period thickness $\Delta z_{cr}$ roughly scales with the phonon mean free path $l$.

Following the above idea of crossover, if the critical period thickness $\Delta z_{cr}$ is characterized with the mean free path $l$, $\Delta z_{cr}$ of an SWNT should be considerably large due to the expected long $l$. In our previous molecular dynamics (MD) study on the length effect of the thermal conductivity of a pure (5, 5) SWNT at room temperature, the ballistic transport of phonons was observed up to at least a few micrometers. Furthermore, the theoretical calculation of Mingo and Broido suggests that the ballistic limit may exceed tens of micrometers. Therefore, when superlattice SWNTs are less than hundreds of nanometers long, which is a realistic length scale in many applications, the entire system is expected to be within the phonon tunneling range and the zone-folding effect would dominate over the TBR effect. On the contrary, we demonstrate that the thermal conductivity of superlattices can take a minimum value even in much shorter system well in the ballistic phonon transport regime.

As simulations of superlattices in three-dimensional materials inevitably suffer from uncertainties caused by simplifications of the boundary scattering and anisotropic effects, SWNTs would be the ultimate case where the theories can be applied without major simplifications due to the quasi-one-dimensional confinement of phonons. The synthesis of SWNTs from $^{13}$C isotopes source has been demonstrated to be possible. With the recent outstanding development of SWNT growth techniques using chemical vapor deposition in well-controlled conditions, it may be possible to experimentally pattern superlattice structures in the near future.

II. MOLECULAR DYNAMICS SIMULATIONS

Molecular dynamics (MD) simulations utilize Brenner potential with a simplified form to express interaction between carbons. This potential function has been reported to be able to describe variety of small hydrocarbons, graphite, and diamond lattices. The formulation of the potential function is based on the covalent-bonding treatment developed by Tersoff. The total potential energy of the system is expressed as

$$E = \sum_i \sum_{j<i} [V_{ij}(r_{ij}) - B_{ij}^* V_A(r_{ij})],$$

where $V_{ij}(r)$ and $V_A(r)$ are repulsive and attractive force terms, which take the Morse type form with a certain cutoff function. $B_{ij}^*$ represents the effect of the bonding condition of the atoms. As for the potential parameters, we employ the set that was shown to reproduce the force constant better (Table 2 in Ref. 19). The velocity Verlet method was adopted to integrate the equation of motion with the time step of 0.5 fs. Simulations were done for armchair SWNTs with chiral index of (5, 5), which gives the radius of about 0.7 nm.

Superlattice structures were constructed by alternately connecting (5, 5) SWNTs with different masses. Figure 1 denotes an isotope superlattice with cells of $^{12}$C and $^{13}$C with a certain period thickness $\Delta z$. Simulations for $^{12}$C/$^{24}$C system were also carried out for clearer demonstration of the influence of $\Delta z$ on the heat conduction. Although the former case is more realistic, as shown later, the fluctuations severely influence the statistics even with a considerable number of sampled data. Therefore, the latter case with higher mass ratio serves to highlight the targeted phenomenon by enhancing the signal-to-noise ratio.

The methodology of the thermal conductivity measurement follows our previous reports. The temperature gradient was applied by using the phantom atoms placed on both ends of an SWNT. Once a quasistationary temperature profile is achieved, $\lambda$ can be calculated from the temperature gradient and the heat flux $q$ obtained from energy budgets of phantom atoms using the Fourier’s law, $q = \frac{Q}{A} = -\lambda \partial T/\partial z$. Here, the cross-sectional area $A$ is defined as the area of a hexagon dividing a close-packed bundle of SWNTs $A = \frac{2\sqrt{3}}{b} (d/2 + b/2)^2$, where $b$ is van der Waals thickness 0.337 nm. Although, the definition of $A$ is arguable, the intention here is simply not to exaggerate the value of $\lambda$ by choosing the definition that would result in relatively small $A$.

Simulations were performed with different $\Delta z$ and SWNT length $L$. Before imposing the temperature gradient, the average temperature of the system was set at 300 K with the auxiliary velocity scaling control. Measurements were started when the quasistationary state was achieved by the phantom thermostats maintained at 290 K ($z = -L/2$) and 310 K ($z = L/2$). The measured data were averaged over 10–20 ns. Within the explored parameter space, the time-averaged temperature profile remained quasilinear. As seen in the temperature profiles for $\Delta z = 0.5$ nm [Fig. 2(a)] and 5.0 nm [Fig. 2(b)], the temperature profiles exhibit negligibly small bumps with the length scale corresponding to $\Delta z/2$. Heat fluxes added to and subtracted from the phantom thermostats at the tube ends remained almost constant as those in the simulations of pure $^{12}$C-SWNTs.

III. RESULTS AND DISCUSSIONS

A. Minima in thermal conductivity of $^{12}$C/$^{13}$C superlattices

The result of the simulations for $^{12}$C/$^{13}$C SWNT superlattice at room temperature shows that $\lambda$ takes a minimum value at $\Delta z_{cr} \sim 5$ nm (Fig. 3). The value is apparently much smaller than the expected $l$ of SWNTs. Although, the mean free path of the SWNT superlattice may well be shorter than that of a pure $^{12}$C-SWNT, the reduction of the mean free path
should be limited to at most 50% as estimated by the reduction of the thermal conductivity. Let us now consider the contribution of individual phonons to heat conduction, i.e., \(\lambda_i\), in the regime with dominant influence of TBR \(\Delta z\). We assign phonons into three categories with different contribution to \(\lambda\); (1) fully ballistic phonons \((l_i \geqslant L)\) which have \(\Delta z\)-invariant contribution to the conduction due to the phonon tunneling effect, (2) diffusive-ballistic phonons with \(0 < l_i < L\), whose contribution to \(\lambda\) is \(\Delta z\) dependent, and (3) stationary phonons with negligible contribution to heat conduction due to the small product of group velocity and mean free path, \(v_i l_i\). The phonons in the second category are the only ones with noticeable contribution to the dependence of \(\lambda\) on \(\Delta z\). Therefore, if there is any mean free path that should scale with \(\Delta z_{cr}\) that would be the local mean free path of the diffusive-ballistic phonons. Yet, in MD simulations and also in reality, unlike the simple theoretical models with a single scale of mean free path, \(l\) has a certain distribution and there is no assurance that it takes a simple form. Hence, the precise scale of \(\Delta z_{cr}\) should depend on the distribution function of \(l_i\) weighted by \(C_i v_i\).

Thermal conductivity of an SWNT is known to exhibit distinct length effect, where \(\lambda\) increases with the nanotube length. On varying \(L\) from 50 nm to 101 nm, \(\lambda\) of the pure \(^{12}\)C-SWNT \(\lambda_0\) increases from 313 W/mK to 370 W/mK due to the length effect. Accordingly, \(\lambda\) of \(^{13}\)C/\(^{12}\)C SWNT superlattices increase with \(L\) for the entire range of \(\Delta z\). On the other hand, as seen in Fig. 3, \(\Delta z_{cr}\) appears to be insensitive to the change of \(L\). This is consistent with the above discussion that the fully ballistic phonons added by lengthening the nanotube simply increases the offset \(\Delta z\)-independent thermal conductivity and have minor influence to the dependence of \(\lambda\) on \(\Delta z\).

In our previous MD studies on the length effect of the SWNT thermal conductivity, the heat conduction exhibited a strong one-dimensional nature in the current length regime. Assuming the SWNT superlattice to be a one-dimensional system, the superlattice with the smallest \(\Delta z\) which consists of two monolayer unit cells can be considered as an “alloy.” In this sense, the current results can be understood as a case where the attenuation of the thermal conductivity with superlattice structures beats the alloy limit. However, it should be noted that the minimum thermal conductivity is still slightly higher than \(\lambda/\lambda_0=0.53\) \((\lambda/\lambda_0 =0.07\) in case of \(^{12}\)C/\(^{13}\)C system) obtained by randomly mixing \(^{13}\)C atoms into a pure \(^{12}\)C-SWNT with a number ratio of 50%.

**B. Model superlattice system with \(^{12}\)C/\(^{24}\)C**

As seen in Fig. 3, even by sampling for 10–20 ns (20–40 million time steps), the statistics seriously suffer from fluc...
In order to highlight the influence of superlattice structures and validate the above analyses, simulations were carried out for the $^{12}$C/$^{24}$C system ($L=50$ nm). Consequently, as denoted with asterisks in Fig. 4, data are much smoother than in the previous case. Again, the minimum $\lambda$ is observed but this time the picture is more distinct. Larger mass difference results in smaller average $\lambda$ [at $\lambda_0 = 310$ W/mK] compared with the previous case. Correspondingly, $\Delta z_{cr}$ is reduced to 1.5 nm, which can be understood in terms of the narrowing of the phonon tunneling regime, i.e., the reduction in mean free path of diffusive-ballistic phonons with $0 < z < L$ due to the enhancement of the impact of each lattice interface.

Figure 4 also shows the temperature dependence of $\lambda$. On decreasing the temperature from 300 K to 50 K, mean free paths of the diffusive-ballistic phonons are expected to become longer. An apparent consequence is the increase of $\lambda$ due to the attenuation of the thermal phonon scattering (normal and umklapp scatterings). In the current case, thermal conductivity of a pure $^{12}$C-SWNT increases from $\lambda_0 - 310$ W/mK (300 K) to $\lambda_0 = 1167$ W/mK (50 K). Correspondingly, $\Delta z_{cr}$ is shifted upward on the horizontal axis and as a consequence $\lambda$ decreases monotonically without crossing the crossover limit in the entire range of $\Delta z$. The trend indicates that the TBR effect is small, which can be attributed to dominant ballistic phonon transport at low temperature for the current system size.

### C. Observation of phonon dispersion relations for superlattices

The simulations of the $^{12}$C/$^{24}$C system enable us to clearly observe the zone-folding effect in the dispersion relations. The dispersion relations can be obtained from MD simulations by calculating the two-dimensional (2D) Fourier spectra of the time history of the one-dimensional (1D) velocity field along an SWNT. Here, spectra are presented in terms of the energy density in $(\omega,k)$ space,

$$ E(\omega,k) = \frac{1}{3n} \sum_{a} \sum_{z} \left| \frac{1}{N} \int v_{a}(z,t) \exp(ikz-i\omega t) dt dz \right|^2, $$

where $N$ and $n$ are the number of atoms in the longitudinal ($z$) direction (the number of unit cells in the nanotube) and number of atoms in a unit cell, respectively. The velocity vector is projected to the local cylindrical coordinates $(r,\phi,z)$ denoted by the subscript $a$ in Eq. (2). The energy density was first computed for each directional component then summed to obtain the overall dispersion relations. In Fig. 5, $k$ is normalized with the Brillouin-zone boundary length $\pi/a$. In the current case with an armchair SWNT, a unit cell is an armchair-shaped monolayer and hence $a = \sqrt{3}d_{cc}$, where $d_{cc}$ is the interatomic distance. The data are discrete due to the finite length of the nanotube and the broadening of the spectral peaks indicates the finite temperature effect and phonon scattering. The features of the dispersion relations of a pure $^{12}$C-SWNT obtained from equilibrium MD simulations shown in Fig. 5(a) (Refs. 22–24) agree with the ones of the previous models.\(^{23}\) They also agree well with more recent mechanical model with first-second neighbor directed bonds and radial bond-bending interactions reported by Mahan and Jeon.\(^{25}\)

In Figs. 5(b)–5(d), the dispersion spectra of 50 nm long SWNT isotope superlattices subjected to quasistationary temperature gradients are shown for $\Delta z=0.5$, 1.5, and 5.0 nm, which correspond to 2, 6, and 20 monolayers. Each case represents the regime with dominant zone-folding effect ($\Delta z < \Delta z_{cr}$), crossover ($\Delta z \sim \Delta z_{cr}$), and dominant thermal boundary resistance ($\Delta z > \Delta z_{cr}$). The figures are drawn to provide closeups of the low-frequency and wave vector regime capturing the key phonon branches for heat conduction; LA, TA, and $\Gamma$ together with four low-frequency optical phonon branches ($LO_1$ and $TO_1$). As denoted with dashed lines, the periodicity imposed by the superlattices gives rise to the mini-Brillouin zone width $\pi/\Delta z$ in $k$ space normalized by the width of the original Brillouin-zone boundary of a pure $^{12}$C-SWNT. For cases with $\Delta z=2a=0.5$ nm [Fig. 5(b)] and $\Delta z=6a=1.5$ nm [Fig. 5(c)], the zone-folding effects are visualized as the local symmetries of the branches with respect to the mini-Brillouin zone boundaries. The appearance of the mini band gaps at the zone boundaries\(^{11}\) and the corresponding reduction of the group velocity due to the zone-folding effect can be observed. Note that there are distinct discrete jumps of the phonon branches at the intersections with the mini-Brillouin zone boundaries, and the group velocity is locally reduced in the regime near the zone boundaries. Figure 5(d) depicts that, in case of $\Delta z=20a=5$ nm, each mini band gap becomes smaller, if not diminished, compared with the previous cases, which confirms the switch of the dominant effect from the zone-folding to the thermal boundary resistance. These observations of dispersion rela-
tions agree with Simkin and Mahan,\textsuperscript{10} where the recovery of thermal conductivity in $\frac{\Delta z}{10}\frac{\Delta z}{11022}$ is characterized by the gradual closing of the mini band gaps with increasing $\frac{\Delta z}{10}\frac{\Delta z}{11022}$.

**IV. CONCLUSIONS**

Heat conduction of SWNT isotope superlattices was investigated by means of nonequilibrium molecular dynamics simulations. Superlattice structures were constructed by alternately connecting $\frac{\Delta z}{12}\frac{\Delta z}{20849}$-SWNTs with different masses. At room temperature, implementation of $\frac{\Delta z}{12}\frac{\Delta z}{20850}$ superlattice structures can attenuate the thermal conductivity to approximately 56% of the value of the pure $\frac{\Delta z}{12}\frac{\Delta z}{20850}$-SWNT. By performing the simulations over a range of period thicknesses, the thermal conductivity was found to take a minimum value at a certain critical period thickness, which reflects the crossover of the zone-folding and the thermal boundary resistance effects. As a consequence, the reduction of the thermal conductivity of superlattice structures beats that of the one-dimensional alloy structure, though the reduction of thermal conductivity is less than the value obtained by two-dimensional random isotope mixing. The results demonstrate that the minimum thermal conductivity appears even in the system dominated by ballistic phonon transport, depending on the distribution of diffusive-ballistic phonons whose mean free path $l_i$ is in the range of $0 < l_i < L$. Finally, by calculating the two-dimensional energy density spectra, the zone-folding effects were observed together with the mini band gaps, where the opening of mini band gaps increased with $\frac{\Delta z}{12}\frac{\Delta z}{20850}$.

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