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Atomistic Evidence of Hydrodynamic Heat Transfer in Nanowires

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Abstract

With wave-packet propagation simulations and heat flux estimation via molecular dynamics, we show that the heat flux radial distribution in silicon nanowires can be described by a mesoscopic model, the hydrodynamic heat equation. We observe Poiseuille like heat flux profile, that cannot be described by a simple kinetic model such as the Fuchs-Sondheimer model, in both pristine and core/shell nanowires. The addition of a shell does not change the shape of the radial heat flux distribution, but just modifies the maximum of the heat flux in the center of the nanowire. These results show that there is a heat flux depletion length for pristine or core shell nanowires, 1-2 nm away from the boundary of the crystalline part. The parameters of the mesoscopic model are discussed in terms of microscopic properties, including the phonon mean free path as function of frequency and the partial vibrational density of states in the different regions of the nanowire.

Keywords: hydrodynamic heat transport, Nanowires, Amorphous Shell, Nanocomposites

1. Introduction

Heat transfer at the nanoscale has been originally described using the linearized Peierls Boltzmann (PB) equation [1]. Among the pioneering models, the

early kinetic theory models [2], or the Maxwell-Cattaneo model [3] can be men-
tioned. They overcome some of the shortcomings of the conventional Fourier's
law at the nanoscale, by modeling thermal transport in terms of the heat car-
rier dynamics and of interactions among themselves and with the lattice defects.
Since then, numerous models of thermal transport at the nanoscale have been
developed. These approaches can be separated into two main categories: those
based on a continuum approach such as the Callaway Model [4] or Monte Carlo
approaches [5], and those based on discrete representations such as Lattice Dy-
namics [6] or Atomic Green Functions [7].

Among the continuous medium solutions, a method of interest is the kinetic
collective model [8]. It relies on a solution of the linearized PB equation, to
derive the hydrodynamic heat equation [9]. This mesoscopic transport equation
generalizes Fourier's law by including a non-local and a memory term, and it
reduces to the diffusive description for large length and time scales. The original
derivation was proposed to describe small size and high frequency effects in some
specific materials at low temperatures, where momentum preserving phonon-
phonon collisions dominate [10]. However, it has recently been shown that
this description is also relevant for room temperature applications in general
semiconductors like Si [9, 11]. The derived set of equations can be solved using
the finite element method, to model the thermal relaxation of nanostructured
heat sources on Si substrate [12] or thermoreflectance experiments [13].

However, as the size of the object decreases, the continuous modeling reaches
its limits, and the atomic structure of the material has to be explicitly accounted
for. This is possible with Molecular Dynamics (MD). In MD, the trajectories
and interaction forces of the atoms are explicitly computed. The temperature
is defined via the kinetic energy fluctuation, and the heat flux via the inter-
actions and movement of the particles [14]. This allows the computation of
the thermal conductivity via the fluctuation dissipation theorem in the linear
response approximation [15], or more simply by sustaining a thermal gradient
and computing the resulting flux [16]. Alternatively, the thermal transfer can
also be estimated using wave-packet (WP) simulations [17]. MD has the ad-

35 vantage that it does not rely on the phonon gas model, and the whole phonon
spectrum is taken into account. However, as the electronic interactions are not
explicitly considered, it has to rely on an approximated interaction potential.
Moreover, as it is a fully classical method, its results are only valid above the
Debye temperature. Nevertheless, in this temperature range, this method is
40 able to predict correctly the thermal properties of bulk Si [18].

These thermal transfer models are particularly useful to predict the thermal
properties of the nano-objects developed and increasingly used during the last
half century [19]. Silicon nanowires (NWs) are a good example. They were
first discovered in the 60s, and have been used for many applications in the last
45 twenty years [20]. They are a perfect example of small-scale devices for which
the thermal conductivity depends on the size [21]. The reduction of thermal
conductivity in these structures is often linked to non-local effects and phonon-
boundary scattering [22]. In particular, the thermal conductivity is linked to
surface roughness [23]. Interestingly, it can be influenced by the structuration of
50 the NW, as it has been shown experimentally by etching [24] or via simulations
with faceting [25]. A decrease of the thermal conductivity can be obtained,
either by inducing back-scattering or by promoting diffusive scattering over
reflection [23]. This influence of the boundaries leads to a non-uniform radial
flux distribution, that can be observed in MD or in Monte-Carlo solvers of the
55 PB equation [26, 27].

However, Si NWs are rarely pristine. When exposed to oxygen, an
amorphous oxide layer forms [28], and as phonon transmission at an amor-
phous/crystalline interface is possible [29], this may affect the phonon trans-
port in the NW. Moreover, it has been shown that this interface influences the
60 vibrational properties of the crystal near the interface [30]. Likewise, the in-
terface decreases the ballistic heat transport in the NW [31]. The presence of
the shell affects the heat transfer in the core, which can be seen as an effect
of the interface parallel to the heat flux. Thus, the amorphous shell has to be
considered for the estimation of the thermal properties of the crystalline core.
65 Previous studies have already shown the effect of the amorphous shell struc-

turation. For instance, a structural or geometric modulation of the shell can reduce the thermal conductivity [32, 33] or induce rectification [34]. However, the impact of the amorphous layer on the distribution of flux was less studied. The work of Bejenari et al. on the radial temperature profile in NWs [35] and of
70 Verdier et al. on the impact of the amorphization at constant crystalline layer thickness [36] can nevertheless be noted .

In this work, the radial flux distribution in pristine and core/shell Si NWs is obtained with MD and then analyzed at the mesoscopic level of description using the hydrodynamic heat transport equation. Such comparative studies
75 between phonon hydrodynamic transport and MD have already been carried in telescopic Si NW [27], and used to model second sound in germanium [37]. First, the methods used and the NWs studied will be presented. Then, the radial flux distribution obtained with a wave packet method will be presented. Finally, the steady-state flux distribution obtained in non equilibrium molecular dynamics
80 (NEMD) simulations is analyzed using the hydrodynamic heat equation.

2. Methods

2.1. Modeling Core/Shell Nanowires

The configurations used here are silicon NWs, with a crystalline core radius R_{Cry} of 37.5 or 50 Å, and an eventual amorphous shell of thickness e from 10
85 to 20 Å (see figure 1). Their length is equal to 50 nm. The different dimensions, along with crystalline fraction ratio and the radial discretization parameter dr used for heat flux computation, are listed in table 1. To create the core-shell nanowires, the external cylindrical shell is first cut out of an a-Si bulk block, that will be the amorphous shell surrounding the NW (in blue in figure 1).
90 Then this hollow cylinder is filled with c-Si with the $\langle 100 \rangle$ direction of the crystal lattice in the principal direction of the NW. To simulate an infinite NW, periodic boundary condition are used in the longitudinal direction and fixed ones in the other directions. The a-Si sample used was created in an earlier

work [17]. The original sample is a cube of side 17.5 nm repeated three times to
 95 reach the appropriate length.

Table 1: Dimensions of the different configurations with R_{Cry} the crystalline radius, e the amorphous shell thickness, dr the radial resolution of the flux and N_c/N_t the crystalline fraction ratio.

R_{Cry} (Å)	37.5			50		
e (Å)	0	10	20	0	10	20
dr (Å)	2.5	2.5	2.5	2.5	5	5
N_c/N_t (%)	100	62	43	100	69	51

These nanowires are then relaxed with, first a conjugated gradient energy minimization to reach a local mechanical equilibrium, then an equilibration at 100 K with an isothermal/isobaric simulation for 10 ps, followed by an isothermal equilibration at the same temperature for 10 ps. A time step of 1×10^{-3} ps is
 100 used for the equilibrations. After this annealing, a second conjugated gradient energy minimization is performed. We used a modified SW interatomic potential allowing a more realistic modeling of the interface between c-Si and a-Si in terms of interfacial energy and regular atomic energies inside the two phases [30].

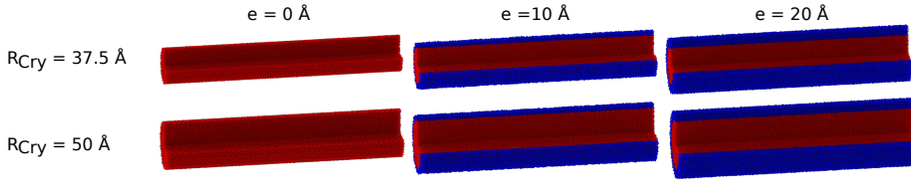


Figure 1: Cross-sectional representation of the different configurations studied, with amorphous atoms in blue and the crystalline atoms in red.

2.2. Kinetic Collective Model

105 The hydrodynamic heat equation relies on a solution of the linearized Peierls Boltzmann equation in a continuous medium to describe heat transport at the microscale [9]. The derivation is based on assuming that, for moderate Knudsen

numbers, the perturbed phonon distribution function is an expansion in terms of the heat flux and its first derivatives in time and space. The solution is obtained for general linearized collision operators and general dispersion relations. Accordingly, the governing equation for the heat flux \mathbf{j} and the temperature T in a crystalline medium is the following:

$$\mathbf{j} + \kappa_{Hyd} \nabla T + \tau \frac{\partial \mathbf{j}}{\partial t} = \ell^2 [\nabla^2 \mathbf{j} + \alpha \nabla (\nabla \cdot \mathbf{j})], \quad (1)$$

with κ_{Hyd} the thermal conductivity, τ the heat flux relaxation time, ℓ the non-local length, α a coefficient [9]. The first two terms of equation (1) correspond to the usual Fourier's law. The last term in the left-hand side introduces the thermal inertial effect, appearing when the timescale is close to the phonon lifetime. The terms in the right-hand side introduce non-local effects that appear when the size of the system is of the order of the mean free path. This last term is analogous to the viscous term in Navier-Stokes equations describing fluid motion. For this reason, equation (1) is called the phonon hydrodynamic equation. If the original derivation was made for low-temperature applications for specific materials like graphene [10, 38], it has been shown that it can be generalized for other materials at room temperature [12, 11]. It is worth mentioning that, in the amorphous shell of the NWs, due to the reduced phonon lifetime, memory and nonlocal effects have been neglected, so the diffusive transport equation (2) is used instead of equation (1):

$$\mathbf{j} + \kappa \nabla T = 0. \quad (2)$$

To predict the thermal response of the system, the transport equation is combined with the energy conservation equation,

$$\nabla \cdot \mathbf{j} = -c_v \frac{\partial T}{\partial t}, \quad (3)$$

with c_v the specific heat. Moreover, appropriate boundary conditions are required. For instance, in the present study, the continuity of the normal heat flux between the crystalline NW and the amorphous shell is imposed. Furthermore, the tangential heat flux \mathbf{j}_t in the crystalline side of the interface is defined

by the slip boundary condition:

$$\mathbf{j}_t = -C\ell\nabla\mathbf{j}_t \cdot \mathbf{n}, \quad (4)$$

with \mathbf{n} the normal-boundary vector pointing towards the amorphous region, and C the slip coefficient weighting the amount of specular and diffusive phonon-boundary reflections [39]. Finally, in the free surfaces of the amorphous shell or the pristine NW, heat flux insulation ($\mathbf{q} \cdot \mathbf{n} = 0$) is imposed.

It is also worth to note that the applicability of the hydrodynamic heat transport model derived in [9] is limited to small enough Knudsen numbers. In the present work, the use of the *ab initio* calculated volumic parameters ℓ and κ_{Hyd} is precluded due to the small NW diameters [39]. However, the model can be still used with effective parameters to interpret the MD simulations as done for previous studies [27].

2.3. Heat Flux in Molecular Dynamics

The goal of this work, is to provide insights about the heat flux radial distribution in core/shell NWs, by comparing with the mesoscopic modeling based on equation (1). This requires the definition of the flux in MD. The heat flux in a volume V can be defined as [14]:

$$J = \frac{1}{V} \left(\sum_i^N E_i \mathbf{v}_i + \bar{\sigma}_i \cdot \mathbf{v}_i \right), \quad (5)$$

with E_i the total energy of the atom i , \mathbf{v}_i the velocity of atom i , $\frac{N}{V}\bar{\sigma}_i$ the stress on atom i and N the number of atoms in the volume. The sum runs on every atom within the volume considered. The virial contribution (part due to the interaction with other particles, as opposed to the energy of the atom itself) $\bar{\sigma}_i$ is computed as follows:

$$\bar{\sigma}_i = - \sum_{k \in K_i} \frac{1}{N_k} \sum_{j \in k} \mathbf{r}_j^k \otimes \mathbf{F}_j^k, \quad (6)$$

with K_i the number of interaction involving the atom i , N_k the number of atom involved in the interaction, \mathbf{r}_j^k the position of the atoms j , \mathbf{F}_j^k the force acting

on atom i due to the interaction with atom j for the k body interaction. This implementation is not exact for many body interaction potentials and correction have been proposed recently [40]. Nevertheless, for silicon the correction does not impact the results in terms of heat flux [41].

160 3. Results

3.1. Ballisticity as a Function of the Frequency

To identify the different energy pathways of the different monochromatic phonons, we have studied the propagation of wave-packets in the nanowires of $R_{Cry} = 37.5 \text{ \AA}$ and $e = 0$ or 10 \AA . These wave-packets are created through the excitation of central layer with a gaussian windowed sinusoidal excitation (see Appendix A.1). The selection of the NWs for this section is motivated by the economy of data, the largest NWs producing larger files.

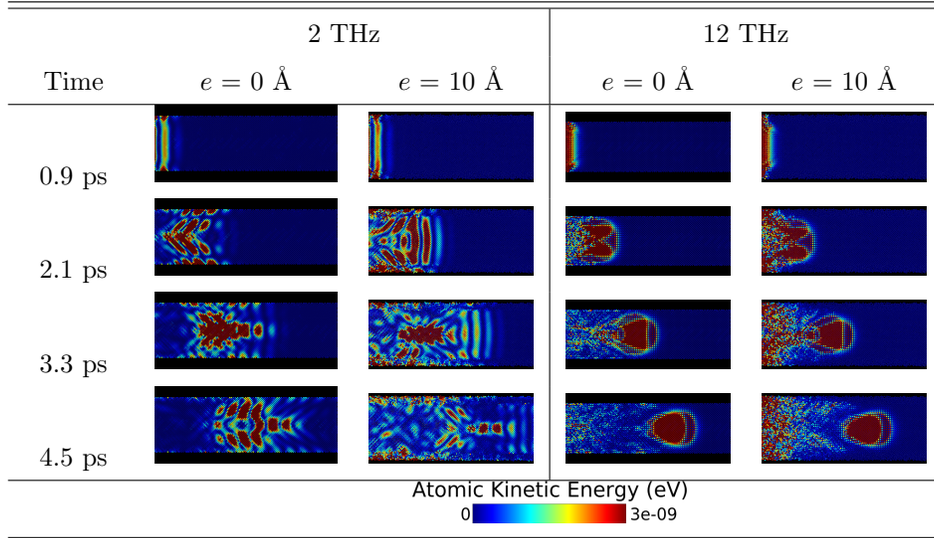


Figure 2: Cross-sectional view of a wave-packet going through the nanowires of $R_{Cry} = 37.5 \text{ \AA}$ and $e = 0$ or 10 \AA . Only the right side of the NW is represented, the propagation on the left side being symmetric. The color scale in the bottom row going from 0 (blue) to $1 \times 10^{-9} \text{ eV}$ (dark red) gives the atomic kinetic energy. Between each line 1.2 ps has elapsed.

The distribution of kinetic energy in a cross-section of the NWs at 2 and

12 THz are depicted in figure 2. At low frequencies, in the first two columns
 170 with or without shell, respectively, the wave-packets initially keep their plane
 wave shape, but are quickly dispersed. The WPs travel at different speeds
 in the different concentric regions of the NWs. This gives rise to interactions
 between the wave in the shell and the wave in the core, as discussed in a previous
 publication [34].

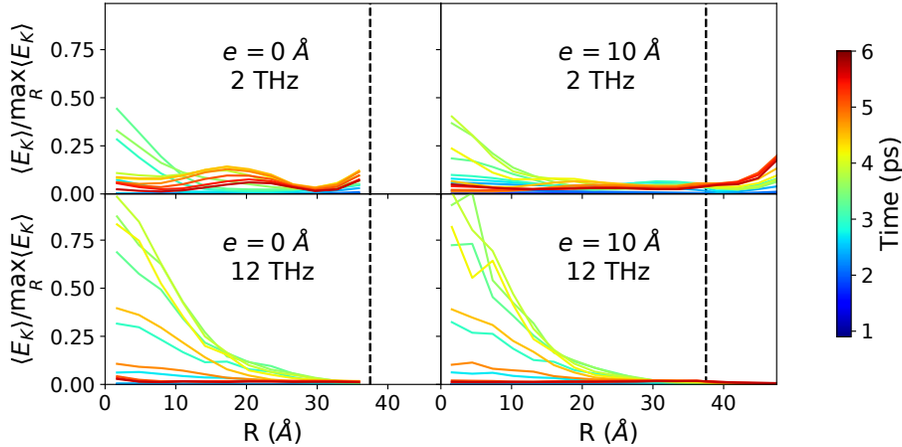


Figure 3: Atomic kinetic energy as a function of the radius averaged on a portion of the NWs 90 to 160 Å away from the excited layer, for different time steps (colors). For the nanowires of $R_{Core} = 37.5$ Å and $e = 0$ Å (first column) or $e = 10$ Å (second column) at 2 THz (first line) and 12 THz (second line). The values are normed by the maximum reached at 12 THz. The vertical black dashed line represents the end of the crystalline core.

175 Moreover, at low frequencies, wave propagation is partly supported by the
 surfaces. This appears in the figure 2, but also in figure 3 that shows that if
 we consider a portion of the NW (here a 70 Å thick slice 90 Å away from the
 excited layer) the energy at the free surface does not stay constant but increases
 with time. This indicates that energy travels at the free surface. It also appears
 180 that more energy travels at the surface in the presence of an amorphous shell.
 Moreover, this figure shows that without shell, a second energy peak appears at
 $R = 20$ Å. This is a consequence of the patterns appearing in the cross-sections.

At 12 THz, the energy propagates differently: it mainly flows in the core of the crystalline region. The profile of the propagating part of the WP is similar with or without shell, as depicted in the bottom panels of figure 3. Little to no energy propagates near the interface/surface, and even less in the amorphous shell. The energy concentration is maximum in the center of the NW and decreases when going toward the surface. This shape indicates a strong effect of the free surface or crystalline/amorphous interface on the energy propagation.

At high frequencies, the main effect of the amorphous shell is the emergence of diffusive transport. This diffusive transport appears as a slowly expanding red patch in the left part of the image in the fourth column of figure 2. This diffusive behavior was originally expected in the amorphous shell [42], more surprisingly, it also appears in the crystalline core when looking at the wave-packet propagation [34].

A more systematic study of the different frequencies shows that, most of the energy is carried by the center of the crystalline core away from the crystalline/amorphous interface or free surface (see Appendix B for more details).

Overall, apart from the complex patterns that appear at 2 THz, the radial distribution of energy during WP propagation is not strongly impacted by the presence of the shell. This short study of the system through WP propagation, shows that even if the free surface seems to transport energy at low frequencies, most of it is transported by the inner part of the NWs away from the interface or free surface. However, the realistic reconstruction of the radial flux distribution from these simulations is not trivial. To access it more easily, NEMD simulations are performed.

3.2. Heat Flux Radial Distribution: Effect of the Shell

To obtain the heat flux as a function of the radius, NEMD simulations are performed, with a thermal gradient of 40 K around 300 K across the NWs (see Appendix A.2). The results, in terms of thermal flux radial distribution, for the nanowires of $R_{Cry} = 37.5 \text{ \AA}$ are reported in figure 4. First, it appears that with or without shell the flux is maximum in the center of the NW, stays constant for

$R < 20 \text{ \AA}$ and then decreases closer to the free surface/interface. This Poiseuille-like heat flux profile is very similar to what was observed by Verdier et al. [26].
 215 In the amorphous shell (browned region), the heat flux does not depend on the radius. There, the reduced phonon lifetime, renders the non-local effects negligible and diffusive transport (equation [2]) is fully recovered. The heat flux profile with or without shell are very similar, the only notable difference is the decrease of the flux in the center of the core in the presence of a shell. An overall
 220 reduction of the flux/thermal conductivity in the crystal due to the addition of an amorphous layer has already been observed for thin films [36].

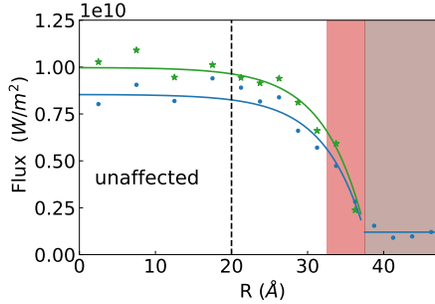


Figure 4: Radial Heat flux in the NW as a function of the radius for $R_{Cryst} = 37.5 \text{ \AA}$ and $e = 0$ (green) or 10 \AA (blue). The dots or stars report the values obtained through the NEMD simulations in the different layers, and the full lines the flux obtained with the hydrodynamic heat transport model (equations [1],[3] and [4]). The dotted line delimits where the shape of the flux distribution is influenced by the boundary (curved profile), and where it is not (flat profile). The brown region delimits the amorphous shell, and the red region shows the crystalline layer where the VDOS is affected due to the presence of the boundary (see figure 5).

We tried to explain the observed profile by changes of the vibrational properties of the NWs due to the free surface or the presence of the amorphous shell. A first explanation for the heat flux depletion near the surface/interface, can
 225 be looked for in the available vibrational modes. For instance, high frequency localized modes could appear in the region where the flux is decreased. Alternatively, a softening could be observed, explaining a local thermal conductivity decrease. To test this hypothesis, the partial vibrational density of states VDOS (see Appendix A.3) in the different layers of the NWs with or without shell

230 are represented in the figure 5. It appears that only the crystalline layer within
 5 \AA of the surface/interface (red dashed line) is affected. This corresponds to
the layer where the structure of the crystal differs the most from bulk c-Si due
to the surface/interface re-structuration or influence of the amorphous shell. In
the inner core, the partial VDOS is very close to the one of c-Si, and in the
235 amorphous shell very close to the one of a-Si. In the end, the perturbations of
the partial VDOS does not extend to the whole region where the flux is reduced
(the red region in the figure 4 does not extend to the dashed line). As a result,
the local flux variations cannot be explained simply by the local modification
of the available modes in the NW core.

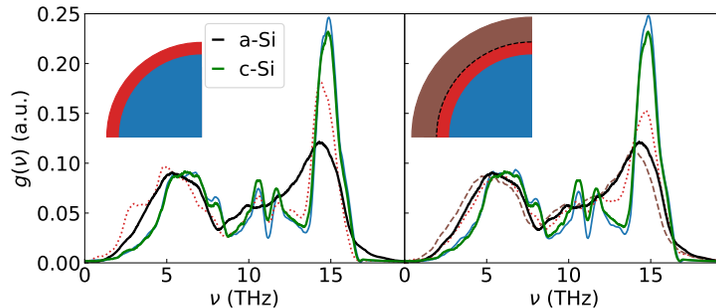


Figure 5: Partial VDOS of the different layers in the nanowires of $R_{Cry} = 37.5 \text{ \AA}$ and $e = 0$ (left panel) or 10 \AA (right panel), the VDOS of bulk a-Si and c-Si are also depicted in, respectively, black and green full lines. The insets give the different colors associated with the different layers of the NW. In the inset, the black dashed line represents the crystalline/amorphous interface.

240 The radial heat flux profiles can be obtained using the hydrodynamic heat
transport model. As explained in the section 2.2, the hydrodynamic heat trans-
port equation is a continuous equation derived from the Boltzmann transport
equation. It adds the influence of memory and non-local effects to the usual dif-
fusible transport mechanism [9]. Here, to reproduce the heat flux profile via the
245 hydrodynamic equations, the same temperature gradient as for the MD simula-
tion is imposed across the NW, and the different parameters of the equations (1)
and (4) are fitted to reproduce the results of NEMD simulations. This is done

using the finite element method, with the software *COMSOL Multiphysics* [39]. Note that, as the steady state is considered reached, the conservation equation (3) is simplified to $\nabla \cdot \mathbf{j} = 0$, and the memory term in the transport equation (1) vanishes ($\frac{d\mathbf{j}}{dt} = 0$). Consequently, the only non-Fourier term that survives is the Laplacian term associated to shear viscosity effects $\ell^2[\nabla^2\mathbf{j}]$. To obtain systematic fits, each parameter is adjusted using different observables obtained in the MD simulations. The thermal conductivity κ_{Hyd} is set first to fit the amount of flux flowing in the central region, where the profile is flat and the viscous effects arising from the boundaries are unnoticeable. The non-local length ℓ is set to fit the size of the region where the flux profile is curved (i.e. the heat flux boundary layer size [39]). Finally, the slip coefficient C is fitted to reproduce the amount of flux flowing at the interface. The solid lines in figure 4, are obtained using this method, for the flux in the crystalline core. In the amorphous shell, the flux does not depend on the radius and is modeled by a Fourier law with $\kappa = 1.5 \text{ W m}^{-1} \text{ K}^{-1}$, which is consistent with the value reported for a-Si using the same inter-atomic potential [43].

The parameters used to obtain the profiles in the crystalline part are reported in the table 2. The easiest parameter to interpret is C , which modifies the flux at the limit of the crystalline core. It depends on the specularly of phonon scattering at the boundary [39]. It appears clearly in the figure 4 that the flux within the last 2.5 \AA crystalline layer before the boundary is more or less the same whether there is an amorphous shell or not. This justifies why the value of C with or without the amorphous shell is the same. On the other hand, the non-local length ℓ characterizes the distance over which the flux is affected by the boundary, and it can be understood as a length over which the flux is self correlated. If assuming the multiscale relaxation time approximation (RTA) to simplify the collision operator, the non-local length can be expressed in terms of the phonon velocities and relaxation times as follows [9]:

$$\ell^2 = \frac{1}{5} \frac{\langle v_{g,\mathbf{q}}^3 / v_{p,\mathbf{q}} \tau_{\mathbf{q}}^2 \rangle}{\langle v_{g,\mathbf{q}} / v_{p,\mathbf{q}} \rangle}. \quad (7)$$

with $v_{g,\mathbf{q}}$ the group velocity, $v_{p,\mathbf{q}}$ the phase velocity, and $\tau_{\mathbf{q}}$ the relaxation time

of the mode \mathbf{q} and the bracket denoting an average weighted by the contribution of each mode \mathbf{q} to the thermal capacity C_v , $\langle x_\lambda \rangle = \int \hbar \omega_\lambda x_\lambda \partial_T f_\lambda^{eq} d\lambda / C_v$. In the present situation, ℓ characterizes how deeply in the core the flux is impacted by the free surface or interface. Again, in figure 4 the flux starts high in the center and is more or less constant until $R = 20 \text{ \AA}$, for both cases. Thus, neither ℓ nor C can explain the heat flux profile modifications appearing with the addition of an amorphous shell. Therefore, the influence of the amorphous shell cannot be directly related with a significant modification of boundary scattering within this model. The decrease of the saturation value is best modeled as a decrease of the effective intrinsic thermal conductivity κ_{Hyd} in equation (1). By using the RTA within the model described by Sendra et. al [9], the thermal conductivity can be expressed as follows:

$$\kappa_{Hyd} = \frac{1}{3} C_v \langle v_{g,\mathbf{q}}^2 \tau_{\mathbf{q}} \rangle, \quad (8)$$

with the same notation as in equation (7).

Table 2: Values of the parameters of equation (1) used to obtain the heat flux profile of the different NWs.

$R_{Cry} \text{ (\AA)}$	$e \text{ (\AA)}$	$\kappa_{Hyd} \text{ (W m}^{-1} \text{ K}^{-1})$	$\ell \text{ (\AA)}$	C	$\kappa_{shell} \text{ (W m}^{-1} \text{ K}^{-1})$
	0	12.5	5	0.3	1.5
37.5	10	10.7	—	—	—
	20	10.7	—	—	—
	0	14	8	—	—
50	10	12.3	—	—	—
	20	12.3	—	—	—

According to expression (8), the variation of κ_{Hyd} might be explained by variations of the group velocity or of the lifetime. However, addition of the amorphous shell does not impact the phonon dispersion in the direction of the propagation (see Appendix C), and thus does not impact the group velocity v_g . An impact over the relaxation times is thus the only remaining explanation of the decrease in thermal conductivity, but the lifetimes also affects the non-local

length, that is constant. However, the lifetimes (or mean free paths $\Lambda_{\mathbf{q}} = \tau_{\mathbf{q}} v_{g,\mathbf{q}}$) are more influenced by the shell for high frequency modes (see Appendix C). As ℓ is less sensitive to the variation of the relaxation time at high frequencies than κ_{Hyd} , it may explain why κ_{Hyd} is impacted by the shell whereas ℓ remains unaffected.

For comparison's sake, the results obtained with MD have also been fitted using an analytical solution of the radial flux distribution in a cylindrical NW derived from the BTE, known as the Fuchs-Sondheimer model. This model has been used in multiple similar study [26]. The analytical solution for a cylinder is given by Dingle and Bragg [44] and reads:

$$\frac{j(r)}{j_0} = \left(1 - \frac{6}{\pi}(1-p) \sum_{\nu=0}^{\infty} p^{\nu} \int_0^{\pi/2} \cos^2 \theta \sin \theta \right) \times \int_0^{\pi/2} \cosh \left(\frac{r \sin \phi}{\Lambda \sin \theta} \right) \exp \left[-\frac{(2\nu+1)\sqrt{R^2-r^2 \cos^2 \phi}}{\Lambda \sin \theta} \right] d\phi d\theta \quad (9)$$

with R the total radius, p the specularly parameter and Λ the MFP. The results obtained with this model in our case are displayed in the figure 6 in grey and purple and compared to the results obtained with MD. To model the flux with equation (9) we used $p = 0$ and two values for Λ . The first value is the average one of c-Si at 300 K, 268 nm [26] and results in the grey curve, the second one is obtained using a least square algorithm to fit the value of Λ to reproduce the NEMD results and results in the purple curve. Finally, we note that increasing the specularly p in expression (9) further decreases the quality of the fits. It appears clearly that neither values of the MFP can reproduce the results obtained by NEMD, in agreement with previous observations for similar NWs [26]. The fit using equation (1) gives more satisfying results, in particular near the interface.

From these observations, one may conclude that an approach that only considers the boundary scattering and a global mean free path cannot reproduce the results obtained with NEMD as well as the hydrodynamic model does.

In summary, it is shown that the heat flux radial distribution in a crystalline

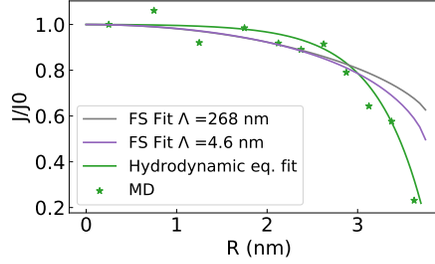


Figure 6: Radial flux distribution for $R_{Cry}=37.5$ without shell, the results from NEMD are displayed as stars, the green line gives the hydrodynamic equations fit, the grey line the fit using the Fuchs-Sondheimer model with the MFP of bulk Si, and the purple line the Fuchs-Sondheimer model using a value of Λ least-square fitted to reproduce NEMD results.

NW is comparable to a Poiseuille flow. Its maximum is in the center of the NW and decreases as it comes closer to the free surface or interface. The flux at the crystalline boundary is similar whether there is an amorphous shell or not. The main effect of the shell is to reduce the saturation flux value in the center of the NW. Such a flux profile can be reproduced with the hydrodynamic heat transport equations (1),(3) and (4), with the effect of shell on the flux in the core reproduced by a decrease of the thermal conductivity parameter κ_{Hyd} in the crystalline core. In the amorphous shell, the flux does not depend on the radius and can be modeled by Fourier's law. Before pushing the analysis further, it is interesting to study the effect of the variation of the shell thickness and crystalline core radius.

3.3. Impact of Diameter and Shell Thickness on Radial Distribution of Heat Flux

In this section, size effects are explored with a study of the impact of the R_{Cry} and e . To this end, both R_{Cry} values and the two shell thicknesses are considered. The flux for the different R_{Cry} are represented in the figure 7, without shell (left panel) or with a 10 or 20 Å shell (right panel). As in figure 4 the dots report the values in the different layers obtained through NEMD simulations, and the full lines the distribution modeled by the hydrodynamic

heat transport equations (1), (3) and (4). The vertical black full and dashed line represent, respectively, surface/interface position for $R_{Cry}=37.5$ and 50 \AA . The profiles are very similar for the two crystalline radii. The main difference is the saturation level that increases upon increasing the crystalline radius. As
 345 observed in figure 4, the flux decreases near the core/shell interface and has the same value in the amorphous shell for the two radii.

The flux distributions for shells of 10 and 20 \AA are displayed in the right panel of figure 7. The reported values are very similar for both shell thicknesses. This shows that the shell thickness does not affect the flux in the crystalline core.
 350 The MD simulated flux values for radial positions below 15 \AA in the case of the small NW and the thicker shell ($R_{Cry} = 37.5 \text{ \AA}$ and $e = 20 \text{ \AA}$ in gray) do not fit this analysis. However, as in the first homoaxial cylinders, the average in equation (5) is performed over less atoms and thus more statistical variations can be expected.

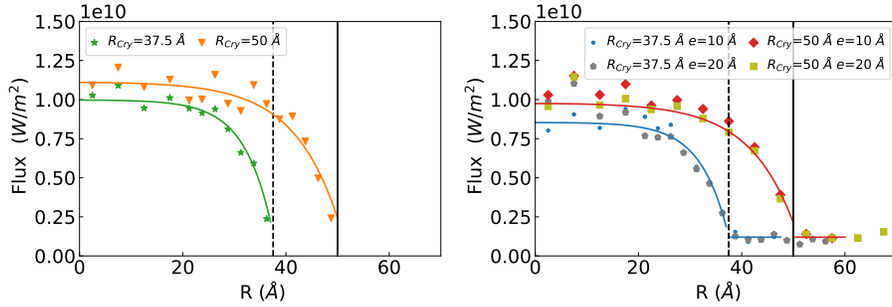


Figure 7: Radial flux distribution for $R_{Cry}=37.5$ and 50 \AA , without shell (left), and with a shell (right). The green, blue, and gray represent respectively the value without shell and with a 10 or 20 \AA thick shell for $R_{Cry}=37.5 \text{ \AA}$. Likewise, the orange, red, and olive represent respectively the value without a shell and with a 10 or 20 \AA thick shell for $R_{Cry}=50 \text{ \AA}$. The full lines represent the flux profile obtained from the hydrodynamic model fits.

355 The parameters used to reproduce the distributions with the hydrodynamic heat transport equations are reported in table 2. It first appears that the same value is used for the slip coefficient C in each case. This could be predicted from the similarity of the flux at the interface in the different cases. Then as

the radius increases, κ_{Hyd} is increased to reach the higher saturation level in
 360 the center of the NW. Again, this value is slightly reduced upon the addition of
 a shell. The non-local length ℓ is also increased from 5 Å to 8 Å when switching
 from $R_{Cry}=37.5$ to 50 Å. A similar effect has already been found for telescopic
 NWs [27]. Again, it is worth noting that both κ_{Hyd} and ℓ in equation (5) are
 linked to the phonon lifetime, that increases as the radius increases, so this can
 365 explain the variations of κ_{Hyd} and ℓ (see appendix Appendix C).

In conclusion, the presence or absence of the shell modifies the radial heat
 flux profile and hence the fitted parameter κ_{Hyd} for the hydrodynamic model
 changes, but modifying the shell thickness has no impact on any parameter
 value. On the other hand, increasing the radius of the core causes both an
 370 increase of the flux in the center of the NW (increase of κ_{Hyd}), and modifies the
 curvature of the heat flux profile (increase of ℓ).

4. Discussion

The radial heat flux distribution in NWs have the shape of a Poiseuille flow
 of fluid in a pipe: the flux is maximum in the center and decreases toward
 375 the interface. For the first time we show that, this shape is not influenced by
 the addition of an amorphous shell, nevertheless the shell impacts the value of
 heat flux in the center of the NWs. The flux decrease appears at a distance of
 20 Å from the free surface or crystalline/amorphous interface. This is similar to
 what was found through MD and Monte-Carlo simulations [26], and with MD
 380 and hydrodynamic heat transport equation without shell [27]. Furthermore, the
 wave-packet propagation hints to a similar heat flux profile.

This flux distribution can be modeled using the hydrodynamic heat equation
 (1) with fitted parameters. Within this model the only parameter that is affected
 by the addition of the amorphous shell is κ_{Hyd} . The increase of R_{Cry} increases
 385 both ℓ and κ_{Hyd} . As the dispersion relations are not affected, neither by R_{Cry}
 nor by the shell presence or thickness (see Appendix C), the variations of
 those parameters are probably due to variation of the MFP/relaxation time, as

discussed in 3.2. However, a decrease in MFP is not the only hypothesis that could explain a decrease of κ_{Hyd} . By definition, the thermal conductivity in equation (1) considers that the behavior is ballistic at the microscale (it is based on the phonon gas picture). However, figure 2 shows that at high frequency, part of the transport is diffusive in the core and in the shell. A partially diffusive transport usually corresponds to an amorphous like thermal conductivity, for which the phonon gas model breaks down [45, 46]. This transition contributes to a decrease of κ_{Hyd} , since the diffusive part becomes more important at the expense of the dominant ballistic part, and manifests itself as decay of the κ_{Hyd} in equation (1).

The lack of impact of the shell thickness on the heat flux radial distribution shape apparently contradicts the results of Shao et al. [47] who reported that the specularly is decreased when the thickness of the shell increases. However, in their definition of specularly they included the diffusion in the amorphous shell, whereas here only the reflections at the crystalline/amorphous interface are considered. This also contradicts the interpretation of the decrease in thermal conductivity in the core-shell NW as an effect of a reduction of the specularly in phonon-boundary collisions due to the presence of the amorphous layer [23]. Indeed, the specularly coefficient in the slip boundary condition (4) characterized from the MD profiles does not depend on the presence of the shell. Therefore, we conclude that the reduction of the global thermal conductivity due to the addition of the shell cannot be explained by reduction of the specularly at the boundary of the crystalline core. Consistently, we did not observe any substantial recrystallisation of the shell nor amorphisation of the crystal that would make the crystalline/amorphous interface rougher than the free surface.

To conclude on the mesoscopic models, it can be noted that the Fuchs-Sondheimer model, which uses a kinetic interpretation of the boundary effects [44], cannot reproduce the results obtained with MD (see figure 6). This suggests that alternative approaches describing the collective behavior of the heat carriers, like the non-local effects included in equation (1), are needed.

The radial energy distribution of the wave-packet in Appendix B can be
 420 compared with the NEMD results in figures 4 and 7. For instance, little energy
 is transported ballistically in the amorphous shell. This is consistent with diffu-
 sive transport in amorphous Si [17]. This diffusive transport explains that the
 heat flux transport in the amorphous shell can be approximated by Fourier’s
 law. Here, we show that the presence of the amorphous shell induces a dif-
 425 fusive transport in the crystalline core as well. Secondly, going closer to the
 interface/surface, there is less energy transmitted. For most of the explored
 frequencies, the highest energy concentration is reached in the center of the
 NW. This energy distribution can be seen as a qualitative measure of the flux
 at different frequencies. The reported energy density, multiplied by the group
 430 velocity, would give an approximate flux at each frequency. However, this ap-
 proach is too crude for a direct heat flux estimation, as the frequencies need
 to be weighted according to the Bose Einstein distribution and the propagation
 direction/polarization considered, as done in previous works [17]. Nevertheless,
 the radial distribution of energy in the WPs indicates that the individual heat
 435 carriers may already favor a Poiseuille-like radial distribution of the heat flux.

The simulations carried out cannot be compared to the continuous model
 within the limit of its validity. As stated in section 2.2, in the original imple-
 mentation of hydrodynamic heat transport equations, ℓ and κ_{Hyd} are estimated
 from *ab initio* simulations on the bulk properties of silicon [9]. This estimation
 440 gives a value of a few hundred of nanometers for ℓ . However, in the present
 case, the diameter of the NW (7.5 nm) is below half this value, so that the *ab*
initio computed parameters are no longer valid [39]. They are replaced by em-
 pirical values, to fit the results obtained with the NEMD method. The resulting
 curves correspond well to the NEMD results, not only in the present study, but
 445 also in a previous work [27]. Even though the *ab initio* calculated parameters
 cannot be used, it is shown here that the phenomena observed in the NEMD
 simulations can be captured with equation (1).

Finally, the determination of thermal conductivity with NEMD shows strong
 size effects [16] so that the obtained radial distribution may depend on the length

450 of the NW. This effect could even impact more the NWs without shell [31]. Even
though, the point of this study is to compare the distribution of heat flux with
or without an amorphous shell at constant length, an impact of the length is
expected since even in NWs of the diameter considered here, phonons with a
large MFP are expected to play an important role on thermal conductivity [48].
455 This importance of long MFP phonons could induce size effects along the NW
longitudinal direction.

5. Conclusion

The radial heat flux distributions in nanowires of diameter 7.5 or 10 nm,
pure pristine or with an amorphous shell, have been studied with three meth-
460 ods: wave-packet propagation, NEMD and the hydrodynamic heat equation.
For all NWs, the radial heat flux profile in the crystalline part is similar: it
looks like a Poiseuille fluid flow. The addition of an amorphous shell decreases
the heat flux in the center of the NW, but the flow at the boundary of the crys-
talline part is not affected by the presence of a shell. To go further, this profile
465 can be reproduced with the equations (1), (3) and (4), using empirical values
for the equation parameters. Doing so, it appears that the addition of the shell
can be reproduced by a decrease in the thermal conductivity used to model the
transport in the crystalline core. The non-Fourier parameters of the equation
do not appear to be impacted by the shell for the geometries considered, and
470 can thus be understood as intrinsic parameters of the crystalline core. As a re-
sult, within this model, the effect of the shell is global, rather than localized at
the interface. Accordingly, the amorphous shell does not impact the specular-
ity of the phonon-boundary collisions. The effective thermal conductivity decrease
may be linked to a partially diffusive heat transport, together with a modifica-
475 tion of the MFP spectra, which motivates frequency dependent analysis of the
thermal conductivity [49].

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485 Appendix A. Methods

Appendix A.1. Wave-packet Propagation

The radial distribution of energy propagation in the NWs can also be studied through the propagation of wave-packets. This can help the representation of the flux and to get more physical insights of what is happening at the interfaces. They can, for example, inform on the shape and pathways taken by individual
490 phonons in the nanostructures [43]. The method used here is very similar to the method used for asymmetric core-shell NWs [34]. An impulsion at a given frequency is imposed on a 2 Å slice in the middle of the NW, and the resulting energy propagation is monitored. The WP is generated through a Gaussian windowed sinusoidal force excitation,
495

$$f = A \sin[2\pi\nu(t - 3\tau)] \exp\left[-\frac{(t - 3\tau)^2}{(2\tau^2)}\right], \quad (\text{A.1})$$

that is imposed to the atoms in the central slice. The amplitude A is $3.773 \times 10^{-4} \text{eV \AA}^{-1}$. The spreading of the Gaussian window τ is selected to be sufficiently small to offer a compromise between spatial extension of the WP compared to the system length and the resolution in the frequency space. The
500 used value is 0.36 ps.

In these simulations, the impulsion is the only source of movement, the initial velocities are set to 0, and the system is at mechanical equilibrium. The simulations are done at constant energy (once the excitation is done), with a

time step of 1×10^{-3} ps. Periodic boundary conditions are used in the direction
 505 of propagation.

Appendix A.2. Thermal Conductivity via Non Equilibrium Molecular Dynamics

To estimate the radial flux distribution in core-shell NWs, the first step is to
 create a thermal gradient across them. This is done by using a hot thermostat
 510 on one side of the NW and a cold one on the other side (as depicted in figure
 A.8a). This set up corresponds to the usual NEMD set up [50].

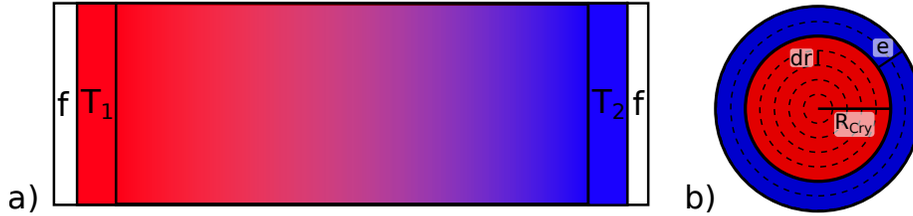


Figure A.8: a) Position of the fixed atoms (f) and hot (T_1) and cold (T_2) thermostat for the
 NEMD simulation, b) schematic representation of the cross-section with R_{Cry} the radius of
 the crystalline core (in red) and e the thickness of the amorphous layer (in blue). The dotted
 lines represent the discretization of the radius for the computation of the flux as a function
 of the radius.

Before establishing the thermal gradient, the NWs are first equilibrated at
 300 K in two steps, a first step with a Nosé-Hoover thermostat and a Nosé-
 Hoover barostat allowing the thermal expansion in the growing direction of the
 515 NWs for 5 ps. In a second step, the boundary conditions are switched from
 periodic to fixed in the growing direction, and the atoms within 10 \AA of the
 extremities are fixed to avoid free surface effects. Once the boundary conditions
 are set, the atoms between the two fixed regions are equilibrated at 300 K for
 50 ps with a Nosé-Hoover thermostat. Only after this step, the atoms from
 520 10 to 20 \AA from the extremities are thermostated at 320 K on one side and at
 280 K on the other. In these thermostats, a simple velocity rescaling is used.
 The trajectories of the atoms are computed using the usual velocity Verlet
 integration scheme, implemented in LAMMPS [51]. The flux is averaged from

1000 ps to 2000 ps, when the steady state is reached. This steady state is defined
 525 here by the stabilization of the temperature profile through the whole NW, and
 the fact that the energy exchanged by the thermostats and the NW increases
 linearly with time. For better statistics, these simulations are repeated 5 times
 for each configuration, with each time a different initial velocity distribution.
 The time step used is of 5×10^{-4} ps.

530 During those simulations, the flux is computed in homoaxial hollow cylinders
 of thickness $dr = 2.5$ or 5 \AA (see figure A.8b). The heat flux in each shell is
 computed using every atom in the shell from 10 \AA away from the cold thermostat
 to 10 \AA away from the hot thermostat. Each studied configuration is listed in
 table 1. For $R_{Cry} = 50 \text{ \AA}$ with $e = 10 \text{ \AA}$ and 20 \AA the dr is larger to decrease
 535 computing time. The flux is computed with the equation (5), sampled every
 0.5 ps, and averaged over the steady state. Equation (5) involves a sum over
 the atoms normalized by a volume, so that in the center of the NW there are
 fewer atoms per shell. This decreased number of atoms increases the variability
 of the computed flux, in particular, for $dr = 2.5 \text{ \AA}$. For this reason, the average
 540 over two homoaxial hollow cylinders is used for radii smaller than 20 \AA .

Table A.3: Global thermal conductivity obtained for the nanowires through the NEMD method using equation 5

R_{Cry} (\AA)	e (\AA)	κ_{MD} ($\text{W m}^{-1} \text{K}^{-1}$)
37.5	0	8.9
	10	5.7
	20	4.3
50	0	11.7
	10	7.5
	20	5.6

In table A.3, the thermal conductivity κ_{MD} extracted from the NEMD simulation is also displayed, in line with what is expected: κ_{MD} decreases with the amorphous shell thickness (e) and increases with R_{Cry} , as reported in previous

studies [36, 32].

545 *Appendix A.3. Vibrational Density of States*

The vibrational density of states (VDOS) are evaluated through the Fourier transform of the velocity auto-correlation function (VACF) [6]. To this end, the system is first equilibrated at 50 K for 100 ps with a Nosé-Hoover thermostat. After this, the thermostats are switched off, and the system evolves at constant energy. Over this 100 ps long simulation, the VACF averaged over the particles 550 belonging to the different concentric homoaxial hollow cylinders of the NWs (see figure A.8b) is computed and recorded. To get the (partial) VDOS, the Fourier transform of the VACF for the different homoaxial cylinders are finally computed and filtered using a polynomial filter.

555 *Appendix A.4. Phonon Dispersion via the Dynamical Structure Factor*

The dynamical structure factor (DSF) is a spatial and temporal Fourier transform of the atomic displacements used to characterize the vibrational properties of a system. It is computed with the same method as in a previous publication [43]. First, the sample is heated to 300 K and equilibrated at this 560 temperature for 50 ps using a Nosé-Hoover thermostat. After this, the atomic trajectories are recorded during a 10 ps long constant energy simulation, the position being recorded every 1×10^{-2} ps. Then, the DSF is computed using the following expression:

$$S(\mathbf{q}, \omega) = \frac{2}{NT} \left| \sum_i^{N_{at}} \exp(-i\mathbf{q} \cdot \mathbf{r}_i) \int_0^\tau \mathbf{u}_i(r_i, t) \cdot \mathbf{m}_\eta \exp(i\omega t) dt \right|^2, \quad (\text{A.2})$$

with \mathbf{q} the wave-vector, \mathbf{u}_i and \mathbf{r}_i the displacement and position of the i^{th} atom, 565 m_η the polarization vector (parallel or perpendicular to \mathbf{q}), T the temperature and N the total number of atoms [46]. The DSF can be obtained for the different vectors of the Brillouin zone. In this study, the focus is laid on the propagation in the axial direction of the NW, that is in the $\langle 100 \rangle$ lattice direction, corresponding to ΓX in the reciprocal space. From the DSF, the phononic dispersion curves 570 can be obtained. For this, the DSF is first filtered through a convolution with

a typical energy resolution curve of line-width 0.33 THz (as suggested by Tlili et al. [17]). Then, for a given wave-vector direction, the dispersion is estimated from the frequency for which the DSF has the highest value for each wave vector within the frequency range of acoustic phonons. Once each wave vector has a value, the curve is filtered using a Savitzky-Golay polynomial filter. The DSF can also be fitted with a damped harmonic oscillator model (DHO) [46],

$$S_{\eta}(q, \nu) = \frac{A}{4\pi^2 ((\nu^2 - \nu_{\eta}^2(q))^2 + \omega^2\Gamma^2)} \quad (\text{A.3})$$

with $\Gamma = 1/\tau$ the inverse lifetime, $\nu(q)$ the phonon dispersion and A the amplitude. This allows the estimation of the lifetime/mean free path.

Appendix B. Frequency Dependence of Radial Energy Distribution

An overview of the behavior at the different frequencies is given in figure B.9. In this figure, the energy distribution profile is given for the timestep for which the highest energy in any homoaxial is reached. This corresponds to the curve of figure 3 at the time step where the highest value is reached locally, for each frequency.

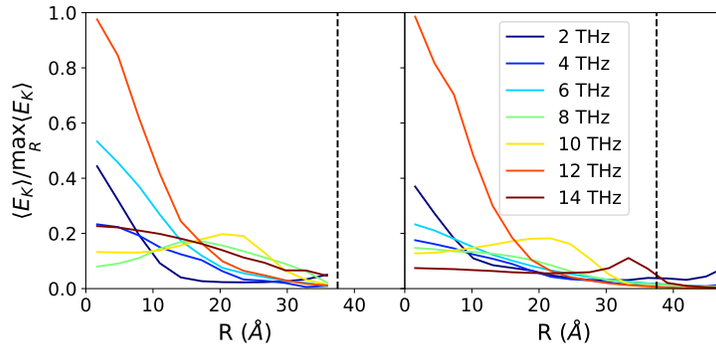


Figure B.9: Atomic kinetic energy as a function of the radius averaged on a portion of the NWs 90 to 160 \AA away from the excited layer, for the time step where the maximal intensity is reached. For the nanowires of $R_{\text{Cry}} = 37.5 \text{\AA}$ and $e = 0$ (left panel) or 10 \AA (right panel). The vertical black dashed line represents the end of the crystalline core.

585 All the frequencies share a common tendency: there is little energy trans-
mitted near the free surface/interface. The only exceptions are at 14 THz where
the maximum is near the interface and at 2 THz with the wave at the surface.
Moreover, the maximal amplitude is not always in the center of the NW: for 8
and 10 THz without shell and for 8 THz with a shell, the maximum is around
590 $R=20 \text{ \AA}$. It is worth noting that the amount of energy injected in the system
depends on the frequency used, so that the quantity of energy at the different
frequencies are not directly comparable.

Appendix C. Dynamical Structure Factor and Mean Free Path

The influence of the shell on the parameters of the hydrodynamic heat trans-
595 port equation could be explained by a modification of the dispersion relation.
Indeed, both κ and ℓ depend on the group velocity (that is, the derivative of
the dispersion relation) and the relaxation time (see equations [7] and [8]). The
DSF for the different configurations are displayed in figure C.10. Even before
the extraction of the dispersion relations, it appears that the main features of
600 the DSF, with or without amorphous shell, are very similar. In both cases, the
acoustic branch appears clearly between 0 and 17 THz. The main difference
between the two is the background noise, appearing in the DSF when there is
an amorphous shell. This corresponds to a-Si modes. It is also to be noted that
a small branch at low frequencies associated to surface modes appears [52]. This
605 is consistent with the observation of the wave at the surface in figure 2. For the
larger diameter (bottom panel), the same observation can be made, though the
background noise is less marked.

The dispersion relations extracted from the DSF, are displayed in figure
C.11. It appears clearly that the different curves superimpose nicely. So that
610 neither the shell nor the radius impact the dispersion relation in the NW for
the propagation in the principal direction, and thus does not impact the group
velocity either.

The remaining microscopic quantity that can be linked to the mesoscopic

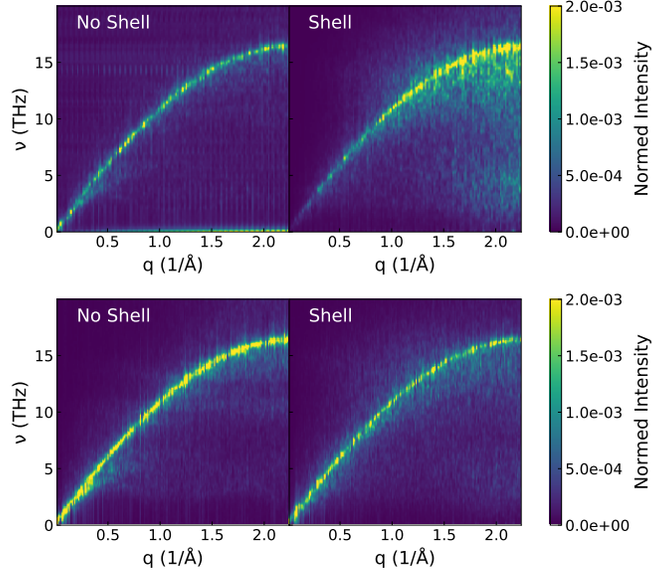


Figure C.10: Dynamical structure factor in the ΓX direction, projected on the longitudinal polarization for the nanowires of $R_{Cry} = 37.5 \text{ \AA}$ (top) and of $R_{Cry} = 50 \text{ \AA}$ (bottom), without (left) or with an amorphous shell of thickness $e = 10 \text{ \AA}$ (right).

transport parameters is the MFP. It can be extracted from the DSF (see section
 615 Appendix A.4) and is given by the enlargement of the DSF around the line of
 maximum intensity. The results are displayed in figure C.11. The MFP of high
 frequencies phonons, above 6 THz, is smaller in the presence of an amorphous
 shell. This variation can be linked to the linewidth increase (and thus MFP
 decrease) due to the amorphous modes visible in figure C.10 for the NW with
 620 an amorphous shell. The decrease of the MFP at high frequencies induced by
 the shell appears clearly for the two radii. However, as the MFP with a shell
 for the two radii are very close, it is not possible to conclude that the increase
 in thermal conductivity linked to a variation of the MFP in this direction. The
 MFPs obtained with this method are notably small compared to the results
 625 obtained for similar NWs using the time domain normal-mode analysis [53]
 but of the order of the results obtained with wave-packet propagation [34].
 However, the large increase in MFP at high frequencies observed with wave-

packet propagation is notably absent when computed with this method.

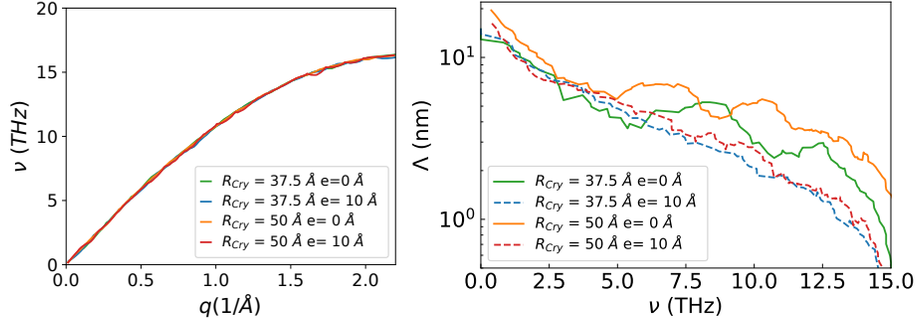


Figure C.11: Dispersion relations extracted from the DSF (right), MFP as extracted using the DHO from the DSF for the nanowires of $R_{Cry} = 37.5\text{-}50 \text{ \AA}$ and $e = 0$ (green-orange) or 10 \AA (blue-red).

This MFP decrease, due to the addition of the amorphous shell, could partially explain the decrease in thermal conductivity. Even if the relaxation time appears in both expressions of ℓ and κ , it is not weighted equivalently in both expressions. Consequently, the MFP variations within a certain range of frequencies may affect the two parameters differently. As ℓ is less sensitive to the variation of the relaxation time at high frequencies than κ , the variation of the MFP as a function of frequency may explain why κ is affected by the shell whereas ℓ is unaffected. However, as the MFP is measured on the whole NW including the amorphous shell the decrease obtained in figure C.11 might be overestimated. Finally, part of the energy could also be supported by diffusive (non-ballistic) phonons.

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