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Phonon thermal conductivity of scandium nitride for thermoelectrics from first-principles calculations and thin-film growth

Sit Kerdsongpanya,^{1,*} Olle Hellman,^{1,2} Bo Sun,³ Yee Kan Koh,³ Jun Lu,¹ Ngo Van Nong,⁴ Sergei I. Simak,¹ Björn Alling,^{1,5} and Per Eklund^{1,†}

¹Department of Physics, Chemistry, and Biology (IFM), Linköping University, SE-581 83 Linköping, Sweden

²Department of Applied Physics and Materials Science, California Institute of Technology, Pasadena, California 91125, USA

³Department of Mechanical Engineering, National University of Singapore, Block EA, 9 Engineering Drive 1, #07-08, 117576 Singapore

⁴Department of Energy Conversion and Storage, Technical University of Denmark, Risø Campus, Frederiksborgvej 399,

Building 779, 4000 Roskilde, Denmark

⁵Max-Planck-Institut für Eisenforschung GmbH, D-40237 Düsseldorf, Germany (Received 2 July 2016; revised manuscript received 6 October 2017; published xxxxx)

The knowledge of lattice thermal conductivity of materials under realistic conditions is vitally important since many modern technologies require either high or low thermal conductivity. Here, we propose a theoretical model for determining lattice thermal conductivity, which takes into account the effect of microstructure. It is based on *ab initio* description that includes the temperature dependence of the interatomic force constants and treats anharmonic lattice vibrations. We choose ScN as a model system, comparing the computational predictions to the experimental data by time-domain thermoreflectance. Our experimental results show a trend of reduction in lattice thermal conductivity with decreasing domain size predicted by the theoretical model. These results suggest a possibility to control thermal conductivity by microstructural tailoring and provide a predictive tool for the effect of the microstructure on the lattice thermal conductivity of materials based on *ab initio* calculations.

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Design of modern materials inevitably requires taking the 24 thermal conductivity into account [1]. For thermoelectric 25 materials a low thermal conductivity is crucial to avoid 26 heat transfer across the legs and therefore loss in energy 27 conversion efficiency [2]. In contrast, electronic components 28 require packaging materials with high thermal conductivity to 29 efficiently dissipate the generated heat [3]. Furthermore, the 30 more complex the devices become, the higher is the demand 31 to control the interplay between microstructure and thermal 32 performance [4,5]. For hard protective wear-resistant coatings, 33 in, e.g., cutting or milling applications, thermal properties 34 are often barely considered when optimizing mechanical and 35 tribological properties, despite the fact that the workpiece is 36 typically subject to temperatures locally exceeding 1100°C 37 in the contact spots [6-8]. The requirements for thermal 38 conductivity are therefore high: the in-plane heat spread within 39 the coating should be as high as possible to ensure uniform 40 heating, while the cross-plane thermal conductivity should 41 be as low as possible to minimize the heat load on the 42 substrate [6-8]. These examples underscore how important it 43 is to understand and be able to tailor thermal conductivity in 44 materials in a broad range of applications. 45

Measurements of thermal conductivity are relatively stan-46 dard for bulk materials but much more challenging for thin 47 48 films and nanoscale materials, still being the topic of active method development [9,10]. There is therefore a substantial 49 need to also develop theoretical methods for predicting or 50 simulating the thermal conductivity of real materials. Recent 51 advances in methodology allow us to predict the thermal 52

*Present address: Department of Materials Science and Engineering, Rensselaer Polytechnic Institute, Troy, New York 12180, USA. [†]Corresponding author: per.eklund@liu.se

conductivity from first principles. In the present paper, 53 we combine state-of-the-art computational techniques with 54 characterization and measurements to determine the effect of 55 microstructure on the thermal conductivity of ScN thin films. 56

When discussing thermal properties from a computational 57 perspective, the lattice dynamical Hamiltonian 58

$$H = U_0 + \sum_i \frac{p_i^2}{2m_i} + \frac{1}{2!} \sum_{ij\alpha\beta} \Phi_{ij}^{\alpha\beta} u_i^{\alpha} u_j^{\beta} + \frac{1}{3!} \sum_{ijk\alpha\beta\gamma} \Phi_{ijk}^{\alpha\beta\gamma} u_i^{\alpha} u_j^{\beta} u_k^{\gamma} + \dots,$$
(1)

is the starting point. It relates the energy of the lattice 59 to displacements (u) of atoms from equilibrium, with a 60 proportionality constant (Φ) for each pair, triplet, quartet, and 61 so on, where *ijk* are indices of atoms and $\alpha\beta\gamma$ are Cartesian 62 indices. Traditionally, these interatomic force constants Φ_{63} are determined by the Taylor expansion of the zero-Kelvin 64 energy around the equilibrium positions, assuming that this 65 expansion is also valid at high temperatures. In this paper we 66 use a recently developed technique, the temperature-dependent 67 effective potential (TDEP) [11–13], where Φ 's are extracted 68 from the finite-temperature *ab initio* molecular dynamics. 69 This provides their intrinsic temperature dependence and the 70 ability to model phenomena inaccessible by traditional means 71 [14,15].

To address these general research questions, we choose 73 scandium nitride as a model system. There is an increasing 74 interest in ScN for a wide range of applications [16], such 75 as a dislocation-reducing buffer for group III nitrides [17] or 76 as a potential thermoelectric material because of its relatively 77 large Seebeck coefficient S accompanied by low-resistivity 78 ρ that results in a high thermoelectric power factor (S^2/ρ) 79 of about $2.5-3.3 \times 10^{-3} \,\mathrm{W \, m^{-1} \, K^{-2}}$ [18–21]. ScN can also 80

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⁸¹ be made p type by doping on Sc sites [22,23]. The former application requires a high thermal conductivity, while the 82 latter requires it to be low. With an indirect gap of 0.9 eV, 83 the thermal conductivity of ScN will be dominated by the 84 lattice contribution [24–27]. In an ideal harmonic material, 85 nothing prevents phonons from carrying heat. In practice, 86 scattering from impurities or isotopes, grain boundaries, and 87 the intrinsic anharmonicity result in a finite phonon lifetime. 88 Sc is isotopically pure. A large single crystal of pure ScN 89 would leave the intrinsic anharmonicity as the sole scattering 90 mechanism. Any alloying dopant would decrease thermal 91 conductivity more than in a material with natural isotopic 92 defects, as the majority of heat is carried by acoustic phonons, 93 rendering the N isotopic distribution less important. This is 94 therefore an opportunity for tailoring the thermal conductivity, 95 by controlling the microstructure. 96

The temperature-dependent phonon thermal conductivity of 97 ScN was obtained with ab initio Born-Oppenheimer molecular 98 dynamics (AIMD). All calculations in this work were carried 99 100 out using the projector augmented wave method [28] as implemented in the Vienna Ab initio Simulation Package 101 (VASP) [29–32]. The AIMD temperature was set to 400, 800, 102 1200, and 1600 K and the volumes to 83.74, 88.12, 92.65, 103 97.34, and 103.82 Å^3 per unit cell for 20 calculations in 104 total, where each calculation was run for 20000 time steps 105 of 1 fs each with a $5 \times 5 \times 5$ supercell (250 atoms in total). 106 Temperature was controlled with a Nosé-Hoover thermostat 107 [33,34]. The exchange correlation was approximated with the 108 AM05 approximation [35,36] and we used the Γ point for 109 the Brillouin-zone integration along with a plane-wave cutoff 110 of 400 eV. The AIMD data were postprocessed using the 111 TDEP [11-13] and the free Gibbs free-energy surface was 112 constructed. From that we extracted volume as a function of 113 temperature, and interpolated the interatomic force constants 114 to the volumes corresponding to zero pressure at the respective 115 temperatures. At these volumes we used the full solution 116 to the linearized phonon Boltzmann equation [37,38] and 117 calculated the lattice contribution to the thermal conductivity 118 on a $31 \times 31 \times 31$ *q*-point grid. 119

ScN thin films were grown on one-side-polished sapphire 120 Al₂O₃(0001) (dimensions $10 \times 10 \times 0.5 \text{ mm}^3$) by reactive 121 dc magnetron sputtering in a high-vacuum chamber with a 122 base pressure of 2.7×10^{-5} Pa. The system is described in 123 detail elsewhere [39]. The Sc target was a 7.62-cm-diameter 124 disk shape, with 99.95% purity with fluorine as a common 125 impurity. Prior to deposition, the substrates were degreased 126 in an ultrasonic bath with trichloroethylene, acetone, and 127 isopropanol for 5 min each and subsequently blown dry with 128 N₂. The Sc target was operated in dc mode (power regulated) at 129 a power of 120 W. The substrate was rotated during deposition 130 in order to obtain uniform films. A substrate bias of -30 V bias 131 was applied to the substrate during deposition. The depositions 132 were performed in Ar/N_2 atmosphere at the total gas pressure 133 of 0.67 Pa with 0.27 Pa Ar and 0.4 Pa N₂. Before deposition, the 134 substrates were heated in vacuum to the deposition temperature 135 (for 1 h for temperature stabilization and degassing). The 136 deposition temperatures (T_d) were 400, 500, 600, and 700 °C. 137 The crystal structure and average characteristic x-ray 138 scattering domain sizes of as-deposited films were determined 139

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by x-ray diffraction (XRD) with standard θ -2 θ scans using 140 CuKa radiation in a Philips PW 1820 diffractometer. Cross-141 section transmission electron microscopy (TEM) and plan-142 view scanning electron microscopy (SEM) were performed on all samples to investigate film microstructure and morphology. 144 For cross-section TEM, the samples were prepared by gluing 145 two pieces of the sample face to face into a Ti grid, 146 then polishing down to 50- μ m thickness. Ion milling was 147 performed in a Gatan Precision Ion Polishing System at 148 Ar⁺ energy of 5 keV and a gun angle of 5° , with a final 149 polishing step with 2-keV Ar⁺ energy and angle of 2°. TEM 150 characterization was performed using a Tecnai G2 TF20UT 151 with a field-emission gun. A ULVAC-RIKO ZEM3 system 152 was used to measure the Seebeck coefficient and electrical 153 resistivity of the films simultaneously in a low-pressure helium 154 atmosphere from room temperature. The error bar of these 155 measurements is within 7%, and the substrate contribution to 156 the Seebeck coefficient and electrical resistivity is negligible. 157

Thermal conductivities of ScN samples were measured 158 using time-domain thermoreflectance (TDTR). We only briefly 159 describe the measurements here; details of the TDTR setup and 160 implementation can be found in Ref. [40]. Before conducting 161 TDTR measurements, the samples were coated with an \sim 80-162 nm-thick Al as transducer. In TDTR measurements, laser 163 pulses from an ultrafast laser are split into a pump beam 164 and a probe beam. The pump beam is modulated and heats 165 the sample periodically, creating a temperature oscillation 166 in the sample. The probe beam monitors the temperature 167 oscillation at sample surface via thermoreflectance (i.e., 168 change of reflectance with temperature). Since the induced temperature oscillation depends on the thermal properties of 170 the sample, TDTR is an accurate approach to measure the 171 thermal conductance of interface and thermal conductivity of 172 nanostructures [41-43]. In the measurements of ScN samples, 173 we use a $1/e^2$ laser radius of 10 μ m, a modulation frequency of 174 10 MHz, and total laser power of \sim 40 mW to limit the steadystate temperature rise to < 10 K. The thermal conductivities 176 of ScN and thermal conductance of Al/ScN are then derived 177 by comparing the TDTR measurements to the calculations of 178 the thermal model [44]. In the analysis, the thickness of Al 179 transducer is determined by picosecond acoustics [45] while 180 the heat capacity of ScN is from the literature value of bulk 181 ScN [46]. The measurement uncertainties are estimated to be 182 $\sim 6\%$ for all measured thermal conductivities. 183

Figure 1 shows the calculated dynamical structure factor S(q, E) of ScN at room temperature, showing signs of strongly anharmonic behavior. There is a faint satellite peak from the transverse optical (TO) mode around the Γ point, owing to the many available scattering channels between the acoustic and optical branches in this region. This is reminiscent of PbTe and SnTe [15]. While the zone center shows deviations from the assumption that the line shape is Lorentzian (see Supplemental Material [47]), the TO phonons carry little heat and we can therefore assume that the phonon Boltzmann equation is still valid.

At room temperature, using the Boltzmann transport equation for phonons, we predict the lattice contribution to the thermal conductivity to be $20 \text{ Wm}^{-1} \text{ K}^{-1}$. This is assuming an infinite perfect crystal. This should be compared to previously reported experimental values of $8-12 \text{ Wm}^{-1} \text{ K}^{-1}$ ign

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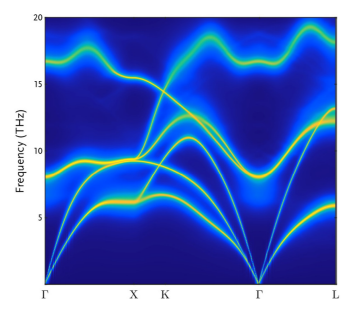


FIG. 1. Dynamical structure factor S(q, E) for ScN at room temperature from first-principles calculations.

[19,48]. The reason for these differences is nonideal effects
of microstructure (and impurities), i.e., that real ScN is not a
perfect crystal. The present synthesis of ScN thin films allows
a systematic, and if not quantitative at least semiquantitative,
correlation between microstructure and other nonideal effects
on thermal conductivity and theoretical calculations of the
same effects.

²⁰⁷ To simulate the microstructural effects, we modify the ²⁰⁸ expression used for the thermal conductivity from

$$\kappa_{\alpha\alpha} = \frac{1}{V} \sum_{\mathbf{q}s} C_{\mathbf{q}s} v_{\alpha\mathbf{q}s}^2 \tau_{\alpha\mathbf{q}s}, \qquad (2)$$

where V is the volume, $v_{\alpha qs}$ is the mode phonon velocity, $\tau_{\alpha qs}$ 209 is the mode phonon lifetime corresponding to wave vector q210 in branch s (with $v_{\alpha qs}$ and $\tau_{\alpha qs}$ determined by an iterative 211 solution to the phonon Boltzmann equation [33,34] and α 212 the Cartesian index), and $C_{qs} = \hbar \omega_{qs} \partial n_{qs}^0 / \partial T$ is the mode specific heat, where \hbar is the reduced Planck's constant $(h/2\pi)$, 213 214 ω_{qs} is the phonon frequency, and n_{qs}^0 is the Bose-Einstein distribution function. By adding the Heaviside step function 215 216 $[\Theta(l)]$ as a function of mean scattering domain size l, the 217 thermal conductivity including the domain-size effect is 218

$$\kappa_{\alpha\alpha}^{\rm acc}(l) = \frac{1}{V} \sum_{\mathbf{q}s} C_{\mathbf{q}s} \upsilon_{\alpha\mathbf{q}s}^2 \tau_{\alpha\mathbf{q}s} \Theta(l - |\upsilon_{\mathbf{q}s}|\tau_{\alpha\mathbf{q}s}).$$
(3)

This means that to simulate a certain mean domain size, 219 we only include those phonons whose mean-free path is 220 small enough to fit inside the domain. The results can be 221 seen in Fig. 2, together with the experimental values. The 222 model here differs from other approaches that model interfaces 223 [49–51], phenomenologically compute the phonon lifetime 224 [52], or directly include boundary scattering in the Boltzmann 225 transport equation [53]. 226

227 XRD results and electron microscopy images of all samples 228 (Figs. 3 and 4) show highly textured polycrystalline ScN films 229 where films were grown in the [111] direction on Al_2O_3 (0001).

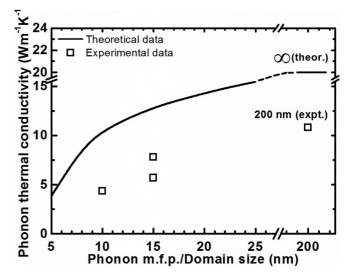


FIG. 2. Theoretical and experimental values of the phonon thermal conductivity of ScN (assuming defect-free crystal) as a function of phonon mean-free path or characteristic x-ray scattering domain size.

In addition, the sample grown at deposition temperature (T_d) ²³⁰ of 700 °C exhibits epitaxial growth, which can be seen from ²³¹ TEM [Fig. 4(f) and Supplemental Material [47]). Samples ²³² grown at $T_d = 400-600$ °C show columnar polycrystalline ²³³ structure and triangular contrast at the interface between film ²³⁴

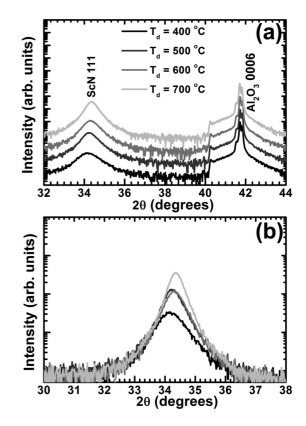


FIG. 3. XRD patterns of ScN films deposited at deposition temperature of $T_d = 400$, 500, 600, and 700 °C on Al₂O₃. (a) Overview and (b) magnified area around the ScN 111 peak.

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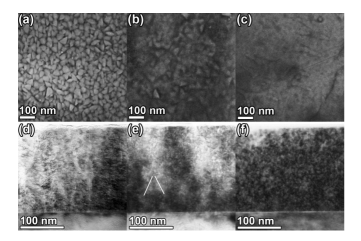


FIG. 4. Electron micrographs of ScN films where (a)–(c) are SEM images and (d)–(f) are cross-section TEM images of the samples that were deposited at deposition temperature, $T_d = 400,600$, and 700 °C, respectively.

²³⁵ and substrate [Figs. 4(d) and 4(e)]. For more information about ²³⁶ samples microstructures see Supplemental Material [47].

From this microstructural information, we see that there 237 are three different types of ScN samples in this study: (1) 238 sample with small domain size, (2) two samples with essen-239 tially the same domain size, and (3) epitaxial thin films, where 240 the domain size is the film thickness. For the polycrystalline 241 films, we can use the characteristic x-ray scattering length 242 in θ -2 θ geometry (corresponding to the peak broadening) in 243 order to estimate the domain size of the film in the out-of-plane 244 direction. The characteristic x-ray scattering length is defined 245 as the size of the (out-of-plane) coherently diffracting domains. 246 Therefore, we can estimate the average out-of-plane domain 247 size using Scherrer's formula [54]. From the XRD results, the 248 2θ full width at half maximum of the ScN 111 peaks that 249 are determined by fitting the curve with Lorentz distribution 250 function are 0.80, 0.54, and 0.56° for the polycrystalline films 251 deposited at 400, 500, and 600 °C, respectively. Thus, the 252 estimates of the average out-of-plane domain size of these ScN 253 films are 10, 15, and 15 nm for $T_{\rm d} = 400$, 500, and 600 °C, 254 respectively. This is consistent with transmission electron 255 microscopy (see Fig. S4 in Supplemental Material [47]). 256

All ScN films were grown to the same thickness of 257 about 200 nm (see Fig. S2 in Supplemental Material [47]). 258 The phonon thermal conductivity of all the samples was 259 determined from $\kappa_{\rm ph} = \kappa_{\rm tot} - \kappa_{\rm e}$, where $\kappa_{\rm tot}$ is the total thermal 260 conductivity and κ_e is the electronic thermal conductivity, 261 which is described as $\kappa_e = LT/\rho$ where L is the Lorenz 262 number $(2.44 \times 10^{-8} \text{ W} \Omega \text{ K}^{-2})$, ρ is the electrical resistivity, 263 and T is the absolute temperature. The electrical resistivity of 264 the polycrystalline ScN films at room temperature is 61.0, 11.0, 265 and 9.8 $\mu\Omega m$ for the films deposited at 400, 500, and 600 $^\circ C,$ 266 respectively. For the epitaxial film, the electrical resistivity 267 is 4.0 $\mu\Omega m$. From the $\kappa_{\rm ph} = \kappa_{\rm tot} - \kappa_{\rm e}$ equation, we can 268 calculate the phonon thermal conductivity from the measured 269 total thermal conductivity at room temperature for all ScN 270 films. The results are shown in Fig. 2. For polycrystalline 271 films, the phonon thermal conductivity of the samples with 272 domain sizes of 10 and 15 nm (two samples) are 5.4, 9.7, 273

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and 7.1 W m⁻¹ K⁻¹, respectively. The origin of the slight ²⁷⁴ difference in phonon thermal conductivity between the two ²⁷⁵ 15-nm-domain-size samples is likely correlated with the fact ²⁷⁶ that the thermal conductance of the Al/ScN interface of ²⁷⁷ the 600 °C sample is also particularly low (see Fig. S3 in ²⁷⁸ Supplemental Material [47]). This indicates that the quality ²⁷⁹ of the sample (in terms of surface oxidation and/or impurity ²⁸⁰ content) is not as good compared to other samples. On the other ²⁸¹ hand, the epitaxial film exhibits a phonon thermal conductivity ²⁸² of 10.5 Wm⁻¹ K⁻¹ which is similar to previous values reported ²⁸³ for epitaxial ScN films [19,55]. ²⁸⁴

Overall, the results show an increase of phonon thermal 285 conductivity with increasing domain size. These results agree 286 well with our calculated phonon thermal conductivity, but with 287 a systematic difference of a factor of ~2. For example, the 288 predicted phonon thermal conductivity of a polycrystalline 289 film with 10-nm domain size is $\sim 10.3 \text{ Wm}^{-1} \text{ K}^{-1}$, while the 290 experimental value is $\sim 5.4 \text{ Wm}^{-1} \text{ K}^{-1}$. The reason for this 291 systematic shift is additional effects, other than domain-size 292 effect, that are not accounted for in the model. Including film 293 thickness and impurity scattering reduces the calculated values 294 close to the experimental ones. Oxygen impurities on N sites 295 (known in ScN films) require high concentrations to affect the 296 thermal conductivity. A concentration of 5 at. % oxygen only 297 reduces thermal conductivity by at most 1% at 300 K. However, 298 since Sc is isotopically pure, any impurities on Sc sites will 299 have a substantial effect on thermal conductivity. Introduction 300 of a fraction of a percent of vacancies, antisite, or substitutional 301 defects (data not shown, as we do not have experimental 302 information on the concentration of Sc-site defects) decreases 303 the phonon lifetimes for the Sc-dominated acoustic branches substantially, lowering the lattice thermal conductivity by as 305 much as 50%. These results and observations validate the 306 model for calculating effects of microstructure on thermal 307 conductivity and enable comparisons of measurements of ther- 308 mal conductivity of real materials with calculations separating 309 different effects. 310

In conclusion, we have theoretically and experimentally 311 investigated phonon thermal conductivity in ScN thin films. 312 Using a theoretical model accounting for microstructural 313 effects on the thermal conductivity, we can semiquantitatively 314 predict the thermal conductivity of ScN. The predictions sys-315 tematically overestimate the thermal conductivity by a factor 316 of ~ 2 , which is due to additional effects, including impurity 317 scattering, which can in principle also be incorporated into the 318 model to improve the quantitative agreement. The dominant 319 effect is domain-size reduction of the thermal conductivity as 320 evidenced by the calculated values of 20 Wm⁻¹ K⁻¹ for an 321 ideal single crystal and 4-13 Wm⁻¹ K⁻¹ for domain sizes of 322 5-20 nm. This large reduction in thermal conductivity with 323 grain size agrees well with our experiments on ScN, and 324 more generally offers an approach for integrated theoretical- 325 experimental design of the thermal conductivity of real 326 materials. 327

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S.K. and O.H. contributed equally to this work.

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