



Applications of Raman spectroscopy in the characterization of nanomaterials

Edited by Paulo T. Araujo and Alexandre R. Paschoal

Imprint

Beilstein Journal of Nanotechnology
www.bjnano.org
ISSN 2190-4286
Email: journals-support@beilstein-institut.de

The *Beilstein Journal of Nanotechnology* is published by the Beilstein-Institut zur Förderung der Chemischen Wissenschaften.

Beilstein-Institut zur Förderung der
Chemischen Wissenschaften
Trakehner Straße 7–9
60487 Frankfurt am Main
Germany
www.beilstein-institut.de

The copyright to this document as a whole, which is published in the *Beilstein Journal of Nanotechnology*, is held by the Beilstein-Institut zur Förderung der Chemischen Wissenschaften. The copyright to the individual articles in this document is held by the respective authors, subject to a Creative Commons Attribution license.



Determining by Raman spectroscopy the average thickness and N -layer-specific surface coverages of MoS_2 thin films with domains much smaller than the laser spot size

Felipe Wasem Klein¹, Jean-Roch Huntzinger¹, Vincent Astié², Damien Voiry³, Romain Parret⁴, Houssine Makhoul¹, Sandrine Juillaguet¹, Jean-Manuel Decams², Sylvie Contreras¹, Périne Landois¹, Ahmed-Azmi Zahab¹, Jean-Louis Sauvajol¹ and Matthieu Paillet^{*1}

Full Research Paper

[Open Access](#)

Address:

¹Laboratoire Charles Coulomb, Université de Montpellier, CNRS, F-34095, Montpellier, France, ²Annealsys, 139 Rue des Walkyries, 34000 Montpellier, France, ³Institut Européen des Membranes, IEM, UMR 5635, Université Montpellier, ENSCM, CNRS, Montpellier, France and ⁴Aix Marseille Université, CNRS, CINAM, UMR 7325, Campus de Luminy, 13288, Marseille, France

Email:

Matthieu Paillet* - matthieu.paillet@umontpellier.fr

* Corresponding author

Keywords:

molybdenum disulfide; number of layers; Raman spectroscopy; thin film; transition metal dichalcogenides

Beilstein J. Nanotechnol. **2024**, *15*, 279–296.
<https://doi.org/10.3762/bjnano.15.26>

Received: 13 November 2023

Accepted: 20 February 2024

Published: 07 March 2024

This article is part of the thematic issue "Applications of Raman spectroscopy in the characterization of nanomaterials".

Guest Editor: P. Araujo



© 2024 Wasem Klein et al.; licensee Beilstein-Institut.
License and terms: see end of document.

Abstract

Raman spectroscopy is a widely used technique to characterize nanomaterials because of its convenience, non-destructiveness, and sensitivity to materials change. The primary purpose of this work is to determine via Raman spectroscopy the average thickness of MoS_2 thin films synthesized by direct liquid injection pulsed-pressure chemical vapor deposition (DLI-PP-CVD). Such samples are constituted of nanoflakes (with a lateral size of typically 50 nm, i.e., well below the laser spot size), with possibly a distribution of thicknesses and twist angles between stacked layers. As an essential preliminary, we first reassess the applicability of different Raman criteria to determine the thicknesses (or layer number, N) of MoS_2 flakes from measurements performed on reference samples, namely well-characterized mechanically exfoliated or standard chemical vapor deposition MoS_2 large flakes deposited on 90 ± 6 nm SiO_2 on Si substrates. Then, we discuss the applicability of the same criteria for significantly different DLI-PP-CVD MoS_2 samples with average thicknesses ranging from sub-monolayer up to three layers. Finally, an original procedure based on the measurement of the intensity of the layer breathing modes is proposed to evaluate the surface coverage for each N (i.e., the ratio between the surface covered by exactly N layers and the total surface) in DLI-PP-CVD MoS_2 samples.

Introduction

The advent of two-dimensional (2D) layered materials beyond graphene has initiated a new field of research [1-3]. In the family of 2D layered structures, transition metal dichalcogenides (TMDs) have attracted considerable attention from academia and regarding potential applications [4-9] because of a number of remarkable properties [10-12]. Particularly, it was found that the properties of layered TMDs drastically change when their thickness is reduced to a monolayer [13,14]. Layered TMD structures have a graphite-like structure with each graphene sheet replaced with an X–M–X or MX₂ triatomic layer, where X is a chalcogen atom (e.g., sulfur, selenium, or tellurium) and M is a transition metal atom (e.g., molybdenum or tungsten) [10].

Among the layered TMD materials, molybdenum disulfide, MoS₂, is of particular interest in optoelectronic applications because of its transition to a direct bandgap semiconductor with very high photoluminescence quantum yield when thinned down to a monolayer [13-17]. Its unique electronic and optical properties could provide an edge in many future applications.

The multilayers MoS₂ structures are of the most common 2Hc type, where atomic layers are arranged in such way that the stacking between two adjacent layers corresponds to a twist angle of $\theta = 60^\circ$, and any Mo atom is sitting on top of two S atoms of the adjacent layers [18,19]. However, during the synthesis process (e.g., chemical vapor deposition (CVD) synthesis) or when using precise transfer or AFM tip manipulation techniques [20], twisted MoS₂ can be formed with two adjacent layers stacked with a relative twist angle (θ) varying from 0 to 60°. Such twisted-layered MoS₂ structures can exhibit a variety of interesting physical properties including unconventional super conductivity [21,22], non-linear optics [23,24], and moiré excitons [25].

Because the properties of MoS₂ flakes are first a function of their thickness, or layer number (N), it is of a primary importance to determine the N of MoS₂ flakes, including twisted MoS₂ flakes and defective MoS₂ flakes, synthesized by different ways. Independently of the structural organization between adjacent layers, a MoS₂ flake is usually named NL -MoS₂, or simply NL , with N being the number of MoS₂ triatomic layers, which defines the thickness of the flake.

Several optical techniques have been developed to identify the N of MoS₂ flakes produced by different methods. Among these techniques, Raman spectroscopy is widely used thanks to its convenience, non-destructiveness, and sensitivity to materials change, including strain, temperature, doping, and defects [26]. Concerning the characterization of MoS₂ flakes, different infor-

mation can be derived from the measurement of the Raman features (frequencies, linewidths, and intensities) of intralayer phonon modes as well as those of the interlayer modes, the so-called layer breathing (LB) modes and shear (S) modes.

Recently, we have developed the reproducible direct growth of wafer-scale MoS₂ thin films on SiO₂/Si substrates by direct liquid injection pulsed-pressure chemical vapor deposition (DLI-PP-CVD) using low-toxicity precursors [27]. Such MoS₂ thin films showed good stoichiometry (Mo/S = 1.94–1.95) and the potential for high photoluminescence quantum yield. However, atomic force microscopy revealed that they are constituted of nanoflakes (with a lateral size of typically 50 nm) with possibly a distribution of thicknesses. Furthermore, depending on the synthesis conditions, the MoS₂ surface coverage can be incomplete, and the thin film average thickness can vary. These samples thus have characteristics, especially thickness inhomogeneities smaller than the laser spot size, that differ from the ones used to establish Raman spectroscopy-based MoS₂ layer counting methods [26,28-33]. In this context, the primary purpose of this work is to develop and validate an approach for determining the average thickness of such sub-laser spot size inhomogeneous MoS₂ thin films using Raman spectroscopy.

First, we reassess here as a ground work the information that can be derived from the Raman spectra of MoS₂ flakes for the evaluation of their thickness, N . Different Raman criteria for the determination of the thicknesses of MoS₂ flakes are first recollected; after the specification of the experimental protocol, domains and limits of application of these criteria are precisely defined from measurements performed on reference samples. These samples are well-characterized, either mechanically exfoliated or standard CVD MoS₂ large flakes deposited on 90 ± 6 nm SiO₂ on Si substrates. Then, we determine which Raman information is relevant to estimate the average thickness of MoS₂ samples produced by the DLI-PP-CVD method, which are constituted of nanoflakes and, thus, significantly different from the reference samples. Finally, an original procedure based on the layer breathing mode intensities is proposed to evaluate the surface coverage for each N , that is, the ratio between the surface covered by exactly N layers and the total surface, in DLI-PP-CVD samples.

Results and Discussion

Experimental procedure

To define a robust experimental Raman protocol to evaluate the thickness of a MoS₂ flake (i.e., its number of layers, N), it is first necessary to specify some parameters that can have a direct influence on the quality and accuracy of the results. The first parameter is the wavelength of the incident laser light used in

the Raman experiments. As it will be detailed in the following, the measurements of the frequencies, linewidths, and intensity of first-order Raman active phonon modes of MoS₂ have to be obtained with good accuracy in order to evaluate the thickness of a MoS₂ flake. These phonons modes are (i) the in-plane phonon mode involving relative motion of Mo and S atoms with E' symmetry for a monolayer (E¹_{2g} for bulk) and (ii) the out-of-plane phonon mode involving only out-of-plane motions of S atoms with A'₁ symmetry for a monolayer (A_{1g} for bulk). These modes are located around 385 and 405 cm⁻¹, respectively, in neutral and defect-free MoS₂ monolayers [33,34]. More precisely, in MoS₂ multilayers, the symmetries of these phonon modes are E' and A'₁ for an odd number of layers, and E_g and A_{1g} for an even number of layers. For simplicity, hereafter when we will discuss the dependence on *N* of the features of these phonon modes, they will be simply referred to as in the bulk, E¹_{2g} and A_{1g}, independently of the number of layers.

A drastic change of the Raman spectra, especially in the frequency range of the A_{1g} and E¹_{2g} modes, occurs when the spectra are excited at an energy close to those of the A and B excitons located around 655 nm (1.89 eV) and 601 nm (2.06 eV), respectively, in MoS₂ monolayers [35,36]. When the incident laser energy is in the range of the A and B exciton energies (the so-called resonance conditions), other bands associated to different second-order processes are observed in the Raman spectra with a strong intensity, their frequencies, widths, and intensity depending on the excitation energy [36]. In addition, resonance conditions alter the symmetry selection rules of phonons of MoS₂ [35]. Some of the second-order bands overlap with the A_{1g} and E¹_{2g} modes, complicating the exact determination of the parameters of these modes recorded under resonance conditions. Furthermore, since the MoS₂ exciton characteristics (energy, width, and spectral weight) can be changed by several factors (e.g., stacking, strain, doping, and defects), the Raman intensities measured with a single laser wavelength close to exciton energies can be affected by external factors and differ for samples elaborated by different methods. For these reasons and in the aim to use Raman spectroscopy to count the number of MoS₂ layers, one must necessarily work under off-resonance conditions, that is, by using incident laser energy far from both exciton resonance energies. In this work, we chose to perform Raman experiments using 532 nm (2.33 eV) laser excitation, because this is sufficiently far from the energy range of A and B excitons [35].

All Raman spectra reported in this paper were recorded on different samples deposited on SiO₂/Si(100) substrate. Hence, the second parameter essential to define is the SiO₂ thickness. Indeed, the multiple reflection interferences that occur in the air/MoS₂/SiO₂/Si structure influence significantly the intensity

of the phonon modes [28,37]. In this work, we chose to focus on substrates with a SiO₂ thickness around 90 nm, which corresponds to the first optimum value for MoS₂ monolayer (*N* = 1) Raman enhancement with a 532 nm excitation energy and also amplifies the signal in the wavelength range of photoluminescence emission (around 650 nm).

The third parameter to define is the power of the 532 nm light, *P*_λ, impinging the sample. Much of the Raman information available to evaluate the thickness of MoS₂ flakes is based on the following parameters: (i) on precise measurements of frequency of the A_{1g} and E¹_{2g} phonon modes of MoS₂. These lead to a precise knowledge of the frequency difference Δω_{A-E}. It was established that Δω_{A-E} depends monotonously on the number of layers, and Δω_{A-E} is largely used as criterion to evaluate the thickness of MoS₂ flakes [26,29,30]; (ii) on the precise evaluation of the integrated intensities of the phonon modes of MoS₂, namely A(A_{1g}) and A(E¹_{2g}), with respect to the integrated intensity of the 521 cm⁻¹ mode from a bare area of the oxidized silicon substrate, A₀(Si), used as an intensity reference [31], or from the silicon substrate underneath the MoS₂ flake, A_{2D}(Si) [28]; (iii) on the precise measurement of the A_{2D}(Si)/A₀(Si) intensity ratio [31]; and (iv) on the measurement of ultralow-frequency modes, the so-called breathing modes and shear modes. The frequencies and the number of LB and S modes allow one to identify the number of layers [32,33] and the presence of twist between adjacent layers from the vanishing of the S modes in twisted MoS₂ flakes [20,38-41].

Then, it is essential to determine the limit value of the laser power so that the above measurements are not affected by laser irradiation. Figure 1 shows the evaluation of the temperature of MoS₂ flakes prepared in different ways and that of the Si substrate as functions of the laser power impinging on the sample through a 100× objective (N.A. 0.9). The power was cycled between ≈5 μW and ≈2 mW. The temperature of MoS₂ flakes is evaluated from the Stokes/anti-Stokes intensity ratio of A_{1g} phonon modes (similar results are obtained using E¹_{2g}) and that of silicon from the Stokes/anti-Stokes intensity ratio of the 521 cm⁻¹ Si mode (see [42] for method details). While the silicon temperature is quasi-insensitive to *P*_λ, the temperature of MoS₂ flakes changes monotonically, reversibly, and quasi-linearly with *P*_λ (see inset of Figure 1). For MoS₂, we found an increase rate of 25–30 °C/mW for monolayers (1L-MoS₂) and 40–45 °C/mW for bilayers (2L-MoS₂). Usual effects of sample heating are the frequency shift of the phonon modes and their concomitant broadening. In Supporting Information File 1, the frequency and the linewidth of the Si mode as functions of the laser power are displayed (Figure S2). These two parameters are found to be insensitive to *P*_λ below 0.5 mW. More intriguing is the evolution of the frequency (Figure 2a) and width (Figure 2b)

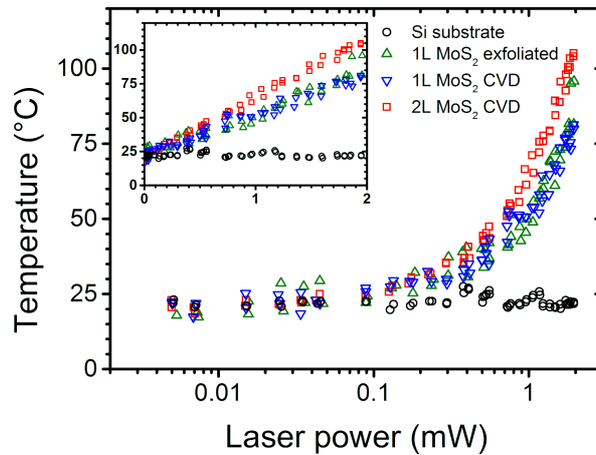


Figure 1: Evolution of the temperature of Si substrate (black circles) and MoS₂ flakes (1L exfoliated: green upward triangles, 1L CVD: blue downward triangles, and 2L CVD: red squares) as functions of the incident laser power in log-scale. The absolute values of the temperature under the laser spot were measured from the variation of the Stokes/AntiStokes ratio of the 521 cm⁻¹ Si mode for the substrate and of the A_{1g} phonon modes for MoS₂ flakes. Inset, same data plotted with a linear laser power scale.

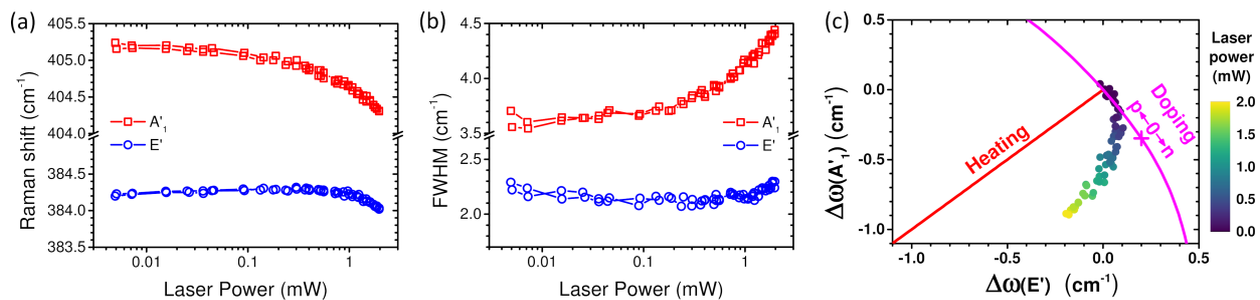


Figure 2: CVD 1L-MoS₂. Evolution of A'₁ (red squares) and E' (blue circles) Raman modes frequencies (a) and full width at half maximum (b) as functions of the incident laser power during a cycle from 5 μW up to 2 mW and back to 5 μW. (c) Evolution of the A'₁ relative shift versus the E' relative shift as a function of laser power. The color code of each point corresponds to the incident laser power as displayed on the color bar. The data are compared to the expected evolutions for heating effect only [43] (red line) and for doping only [44] (magenta line). To enable direct comparison, the strain contribution has been removed, and the corresponding zero doping point is labeled as well as the directions corresponding to p and n doping.

of the phonon modes of 1L-MoS₂ as a function of P_λ . For a simple thermal effect [43] and given the 25–30 °C/mW temperature increase rate determined previously, the frequencies of A'₁ and E' modes should both downshift by 0.3–0.4 cm⁻¹/mW and the width of A'₁ should increase by ≈0.2 cm⁻¹/mW (the width of E' should remain constant) contrary to what is observed in Figure 2a,b. To clarify this point, we present in Figure 2c (filled dots) the relative shift of the frequency of the A'₁ mode versus that of the E' mode measured on CVD 1L-MoS₂ at different laser powers. In the same plot the expected shifts of these modes are reported (i) as functions of a pure thermal effect (Figure 2c, red line) [43] and (ii) as functions of the doping state (Figure 2c, magenta curve) [44]. Clearly the relative shift of the A'₁ mode frequency versus that of the E' mode frequency as a function of P_λ significantly differs from the behavior expected by considering a simple thermal effect. Consequently,

the results reported in Figure 2c clearly evidence photo-doping of 1L-MoS₂ concomitant with a thermal effect, as already observed for MoS₂ on SiO₂/Si [45] as well as for graphene [42]. Furthermore, the evolution of the A'₁ and E' widths with P_λ (Figure 2b), that is, the weak change of the E' width and the significant increase of the A'₁ width concomitant with the A'₁ frequency decrease, support this interpretation [44]. For a laser power smaller than 0.3 mW, photo-doping remains rather low, but it is the dominant contribution to the shift of the modes. Similar results were obtained on other samples including exfoliated 1L-MoS₂. The effects were found irreversible in some cases when P_λ exceeded 1 mW but remained direct always reversible if P_λ was kept below 1 mW.

Based on the above information, all Raman results reported and discussed in this paper were obtained by using a P_λ around

0.1 mW chosen as a good compromise between mitigating laser effects and maintaining measurement efficiency to ensure the accuracy of the Raman criteria discussed in the next part of this paper.

In summary, unless specified otherwise, all Raman spectra reported and discussed in this paper were recorded at an excitation wavelength of 532 nm, with a laser working-power close to 0.1 mW, and using a 100× objective (N.A. 0.9), on MoS₂ flakes or thin films deposited on SiO₂/Si substrate with a SiO₂ thickness of 90 ± 6 nm.

Application of Raman criteria to characterize MoS₂ flakes

In this part, we report and discuss the advantages and limits of some Raman criteria that were found to be efficient to derive the thickness (i.e., the number of layers N) of large MoS₂ flakes prepared by different ways, namely mechanical exfoliation and standard CVD (including twisted CVD 2L-MoS₂). Then, we discuss the application of Raman spectroscopy to characterize samples synthesized by DLI-PP-CVD. In contrast to the first two kinds of MoS₂ samples, the latter are constituted of nanoflakes with possibly a distribution of thicknesses and twist angles between adjacent layers of multilayer domains as well as a higher number of defects.

Exfoliated MoS₂ flakes as reference samples

We performed Raman experiments on mechanically exfoliated MoS₂ [1] that will serve as reference samples. The stacking sequence in exfoliated MoS₂ flakes is of the 2Hc-type [34]. The common feature of all these samples is to have a limited number of defects. Note also that all exfoliated flakes have a lateral size (few micrometers at minimum) significantly larger than the diameter of the laser spot. In such flakes, the exact number of layers, N , is determined by combining optical microscopy, spectral reflectivity, and the measurement of the breathing modes and shear modes in the ultralow frequency (ULF) range of the spectra [32–34].

One of the most popular criteria to determine the number of layers of MoS₂ flakes is the measurement of $\Delta\omega_{A-E}$, that is, the frequency difference between the A_{1g} and E¹_{2g} phonon modes [26,29,30]. Figure 3a shows the dependence of $\Delta\omega_{A-E}$ on the number of layers measured on exfoliated MoS₂ flakes deposited on Si/SiO₂ substrates with four different SiO₂ thicknesses. As previously well documented in the literature, we confirm that $\Delta\omega_{A-E}$ depends monotonously on the number of layers and does not depend on the SiO₂ thickness (Figure 3a). The separation between N and $N + 1$ values are larger than the experimental uncertainties (error bars in the graph) up to $N = 3$. The error bars start to overlap between $N = 4$ and $N = 5$. Comparison with

data from the literature (see inset in Figure 3a) shows that this overlap occurs even between $N = 3$ and $N = 4$ when additional variability due to setup and samples is taken into account. Above $N = 4$, the separation becomes too small compared to the uncertainty. Thus, the measurement of $\Delta\omega_{A-E}$ in exfoliated MoS₂ flakes allows one to evaluate with good accuracy the number of layers for $N \leq 3$. It is then necessary to supplement the $\Delta\omega_{A-E}$ criterion with others to reliably count thick multilayers. In addition, we will establish in the following that this criterion has to be taken with care to derive N in MoS₂ samples other than reference exfoliated MoS₂, because the A_{1g} and E¹_{2g} frequencies, and thus the value of $\Delta\omega_{A-E}$, can be affected by different factors such as stacking order, strain, doping, and defects which can be present in MoS₂ flakes prepared by other ways [44,46–49].

To evaluate the number of layers, we can also use information associated with the integrated intensity of MoS₂ phonon modes. Figure 3b and Figure 3d show, respectively, the dependences of the normalized integrated intensities of the A_{1g} and the E¹_{2g} mode as functions of N for four values of the SiO₂ thickness. For normalization, we use here an external reference, which is a bare Si(111) wafer with only native oxide. In the following, $A(\text{Si}_{111})$ stands for the integrated intensity of the Si(111) 521 cm⁻¹ mode. This reference is preferred to the Si(100) substrate with 90 ± 6 nm SiO₂ to avoid the effects of the SiO₂ thickness variations and crystal orientation. For comparison with other setups or references, the polarization ratio of our setup and the relative values measured on Si(100) with native oxide and 90 nm SiO₂ are given in Supporting Information File 1. As noted by several authors and predicted by the optical interference model, the normalized integrated intensities of MoS₂ modes, namely $A(\text{A}_{1g})/A(\text{Si}_{111})$ and $A(\text{E}_{2g}^1)/A(\text{Si}_{111})$, increase first with N and then decrease showing a maximum for $N = 4–5$ for all SiO₂ thicknesses. Obviously, this non-monotonous dependence prevents using these measurands alone to evaluate the number of layers for $N > 4$. Moreover, a significant dependence of the MoS₂ Raman intensity on the SiO₂ thickness occurs for $N > 2$, pointing out the importance to determine precisely this latter parameter.

Another criterion to derive the thickness of MoS₂ flakes is the $A_{2D}(\text{Si})/A_0(\text{Si})$ intensity ratio [31]. For the evaluation of this ratio, it is of great practical advantage to use the same silicon (the silicon below the oxide, which is Si(100) in the present work) in the measurement of $A_{2D}(\text{Si})$ and $A_0(\text{Si})$. A necessary precaution is that the Si(100) substrate orientation has to be kept the same for both measurements. Another advantage is to give a common origin to the plots of $A_{2D}(\text{Si})/A_0(\text{Si})$ as a function of N ($A_{2D}(\text{Si})/A_0(\text{Si}) = 1$ for $N = 0$) for any SiO₂ thickness.

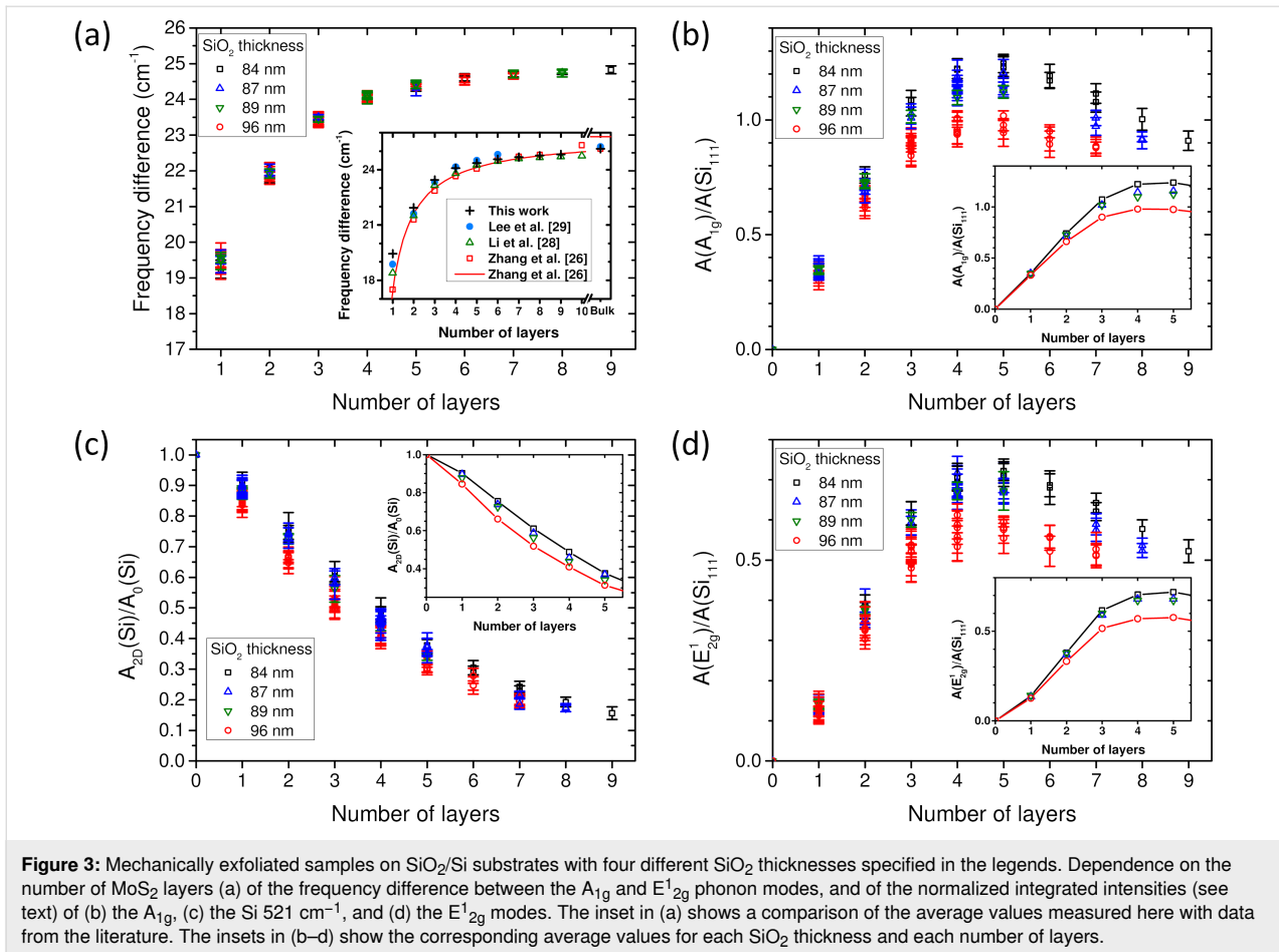


Figure 3c displays the $A_{2D}(Si)/A_0(Si)$ ratio measured on exfoliated MoS₂ flakes deposited on SiO₂/Si(100) substrates with four different SiO₂ thicknesses as a function of N . We confirm the monotonous decrease of the $A_{2D}(Si)/A_0(Si)$ ratio with increasing N [31]. For each N , the value of this ratio depends on the SiO₂ thickness (Figure 3c; black, blue, green, and red symbols correspond to a SiO₂ thickness of 84, 87, 89, and 96 nm, respectively). Despite the monotonous dependence of the $A_{2D}(Si)/A_0(Si)$ ratio, its dependence on SiO₂ thickness combined with experimental errors lead to the conclusion that the measured values for N and $N + 1$ can overlap for any N if the SiO₂ thickness is not known with good accuracy. For a given SiO₂ thickness, the gap between the $A_{2D}(Si)/A_0(Si)$ ratio for N and $N + 1$ is sufficient to ensure a rather good reliability only for $N \leq 5$.

In summary, for exfoliated MoS₂, considering jointly the three Raman criteria (i) value of $\Delta\omega_{A-E}$, (ii) value of the normalized integrated intensities of the A_{1g} and E¹_{2g} modes, and (iii) value of the $A_{2D}(Si)/A_0(Si)$ ratio, one can unambiguously derive the number of layers as long as $N \leq 4$ and the SiO₂ thickness is precisely known. It has also been suggested in the literature

to use the intensity ratio $A(A_{1g})/A_{2D}(Si)$ (or equivalently $A(E^1_{2g})/A_{2D}(Si)$). As it will be discussed in the following, we see two major problems with this approach. The first is the dependence of the Si signal on the crystal orientation and the SiO₂ thickness. The second relates to the fact that using this ratio, rather than using each measurand independently and contrasting them, even if more practical, can hide some information.

Finally, we compare in Supporting Information File 1, Figure S3, the dependence on N of $A(A_{1g})/A(Si_{111})$ and $A_{2D}(Si)/A_0(Si)$ for three SiO₂ thicknesses and two microscope objectives with different numerical apertures, N.A. = 0.9 (blue symbols in Figure S3), and N.A. = 0.5 (red symbols in Figure S3). We observe that the normalized integrated intensity of the A_{1g} mode, $A(A_{1g})/A(Si_{111})$ (Figure S3a–c), is independent of the value of N.A. Concerning the dependence on N of $A_{2D}(Si)/A_0(Si)$ (Figure S3d–f), we found that this ratio is slightly smaller for N.A. = 0.5 than for N.A. = 0.9 and is in a better agreement with the model of Li and coworkers [31] (black solid line in Figure S3d–f). (Note that in this latter work the experimental data on which the model has

been adjusted were recorded using a numerical aperture $N.A. \approx 0.45$).

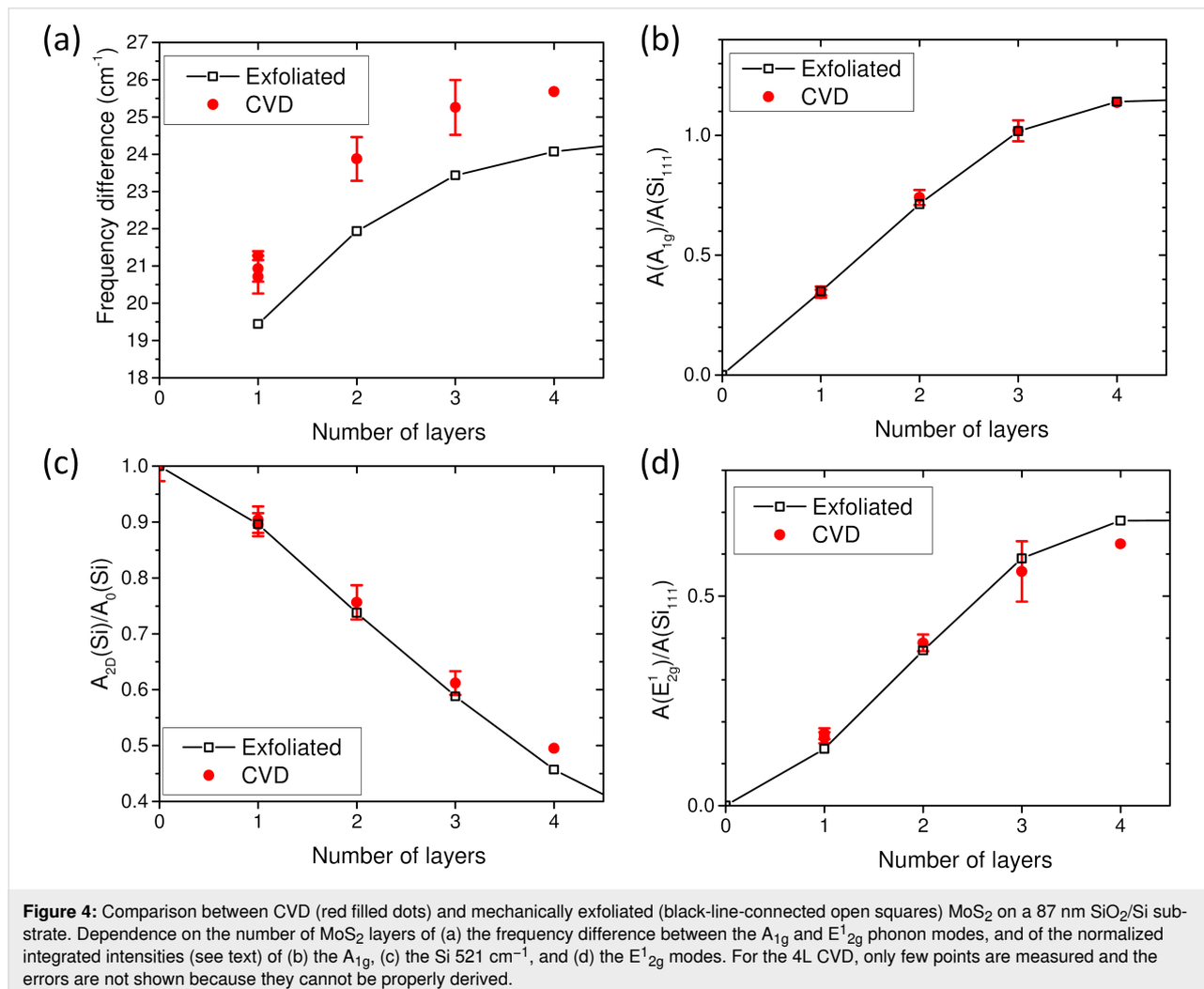
MoS₂ flakes prepared by CVD

In this part, we analyze the pertinence of the previous criteria to derive the thickness of large MoS₂ flakes synthesized by CVD. In a first part, we probe the effectiveness of these criteria to evaluate the thickness of large standard CVD MoS₂ flakes. Such flakes have a limited number of defects and, like in exfoliated MoS₂, the stacking sequence is of the 2Hc type. In the second part, we examine the relevance of these criteria to evaluate the thickness of twisted CVD MoS₂ flakes.

Standard CVD MoS₂ flakes: As derived from the features of the LB and S ultralow-frequency modes, these samples do not show any twist between adjacent layers (presence of S modes for all flakes with $N \geq 2$). The flakes are thus characterized by 2Hc stacking (or close to 2Hc stacking) and a low number of defects, and are named standard CVD MoS₂ flakes. On the

basis of the latter features, the structure of these flakes is close to that of exfoliated MoS₂ flakes. However, the high temperature used in the CVD synthesis and interaction with the substrate can lead to lattice distortion and the presence of vacancies and doping. In the following, we limit our study to a number of layers $N \leq 4$.

Figure 4a compares the values of $\Delta\omega_{A-E}$ measured on exfoliated (Figure 4a, black symbols) and standard CVD MoS₂ flakes (Figure 4a, red symbols) for $N \leq 4$. As previously, the exact number of layers is obtained by combining optical microscopy, spectral reflectivity, and number and frequencies of LB and S modes. For both kinds of MoS₂ flakes, $\Delta\omega_{A-E}$ increases monotonously with N , but for the same N , the values of $\Delta\omega_{A-E}$ are systematically larger in standard CVD MoS₂ flakes. We attribute this discrepancy mainly to a difference of strain states between the two kinds of samples. Exfoliated samples are mostly found with low or slight compressive strain, while CVD samples are under tension. Other factors such as doping, defects,



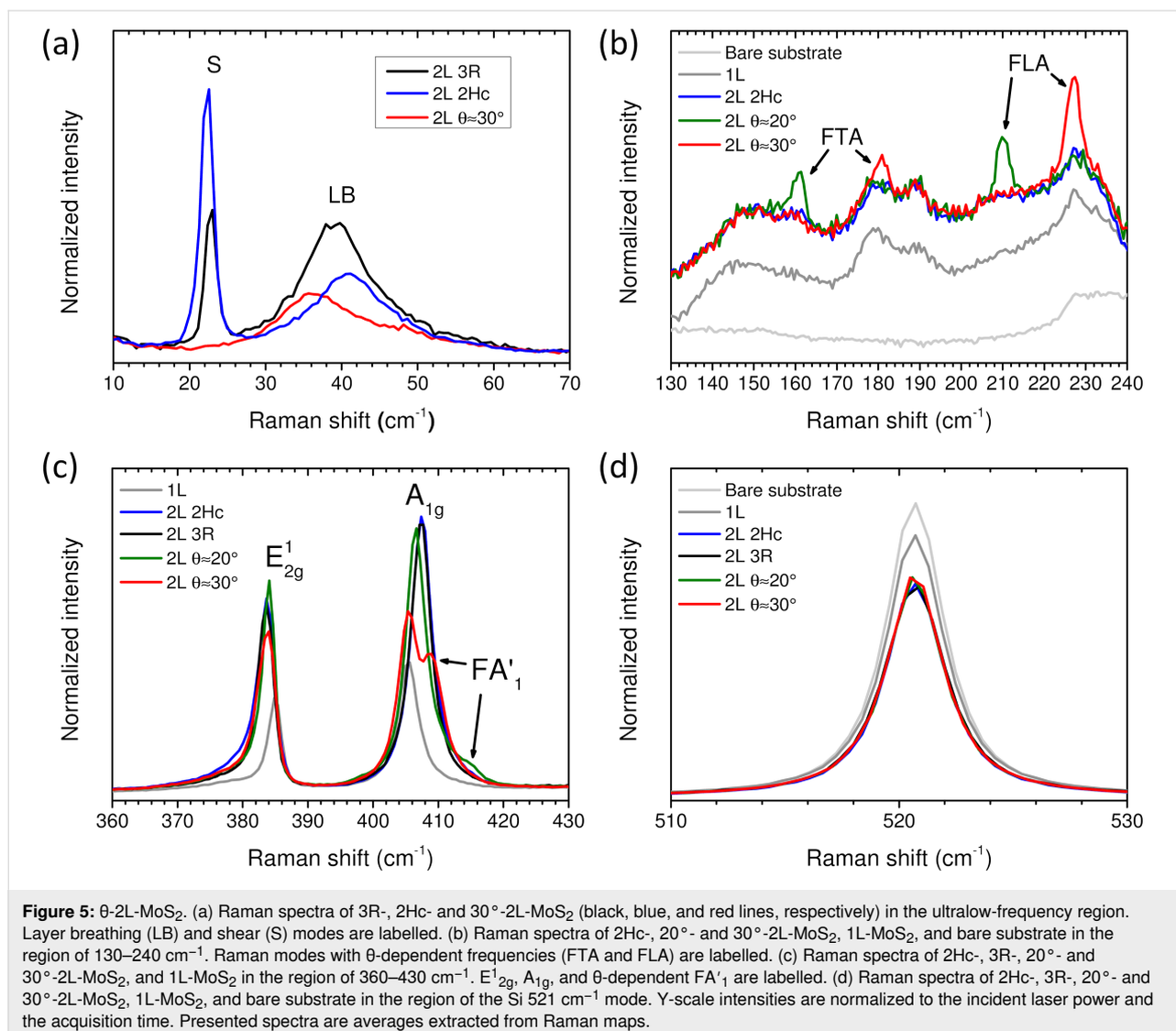
or stacking were shown to lead to large changes of $\Delta\omega_{A-E}$ [44,46,49]. In summary, the value of $\Delta\omega_{A-E}$ is clearly and significantly sample-dependent. Consequently, $\Delta\omega_{A-E}$ cannot be considered as a definitive criterion to derive the number of layers in any of MoS₂ flakes prepared in different ways. In other words, one cannot define a single master curve, $\Delta\omega_{A-E}$ vs N , which would be valid for all the MoS₂ flakes independently of their preparation method or environment.

The dependencies on N of the normalized integrated intensities of A_{1g} and E^1_{2g} modes and the $A_{2D}(Si)/A_0(Si)$ ratio measured on standard CVD flakes are compared with the average values of exfoliated samples with the same substrate SiO₂ thickness (Figure 4b–d). In contrast to $\Delta\omega_{A-E}$, the N dependencies of these intensities are very close in exfoliated and standard CVD MoS₂ flakes. Only $A_{2D}(Si)/A_0(Si)$ and the normalized integrated intensity of the E^1_{2g} phonon modes slightly differ for

$N = 4$. However, this may be due to the fact that the statistics is rather poor on this measurement, because this flake is rather small compared to the others. With regards to these results, these measurands give important information to evaluate the number of layers of 2Hc-stacked MoS₂ flakes independently of the elaboration procedure as long as $N \leq 4$.

Twisted CVD MoS₂ flakes: Other interesting samples are large CVD MoS₂ flakes that present a twist angle, θ , between adjacent layers. We exemplify here the complexity to characterize such samples from the previous Raman criteria with the case of twisted MoS₂ bilayers. The identification of the bilayer character of the investigated flakes was unambiguously obtained independently from spectral reflectivity and optical contrast.

Figure 5a shows the low-frequency range of spectra recorded on three types of MoS₂ bilayer (named 2L-MoS₂ in the following),



namely a bilayer with $\theta \approx 30^\circ$ (this sample belongs to the so-called twisted-bilayer family for which $0 < \theta < 60^\circ$ and is named in the following as θ -2L-MoS₂), the so-called 2Hc-2L-MoS₂ and the so-called 3R-2L-MoS₂. In the latter structure, the stacking between two adjacent layers corresponds to a twist angle of $\theta = 0^\circ$, and it is such that the S atoms of the top monolayer are superimposed on the Mo atoms of the bottom monolayer, and the Mo atoms of the top monolayer are above the hexagon centers of the bottom monolayer [50]. The spectrum in the low-frequency range is dominated by the contributions of the LB and S modes, the frequencies of these modes depending on the twist angle [20,39]. The LB mode emerges in the Raman spectra of all 2L-MoS₂ samples. In line with previous results [20], the Raman shift of the peak position of the LB mode in 3R-2L-MoS₂ is smaller than that of the 2Hc-2L-MoS₂, and the LB mode Raman shift in 30°-2L-MoS₂ is even smaller. Also, in agreement with the literature [20,38–41], the S mode vanishes in 30°-2L-MoS₂.

In Figure 5b, the 130–240 cm⁻¹ range of the Raman spectra recorded on monolayer, 2Hc-2L-MoS₂, and two θ -2L-MoS₂ is displayed. As well documented in the literature, this frequency range is dominated by the contributions of second-order Raman processes [20,26]. The general profile of the spectra is similar in 1L-MoS₂, 2Hc-2L-MoS₂, and θ -2L-MoS₂ with the exception that in the latter flakes, new bands, named FLA and FTA, are superimposed to the second-order Raman spectra. The FLA and FTA modes in θ -2L-MoS₂ are attributed, respectively, to folded longitudinal acoustic phonons and folded transverse acoustic phonons of the monolayer due to the presence of a moiré superlattice [20]. As shown in the literature [20], the frequencies of these modes depend on the twist angle (see Figure 2e in [20]). Unfortunately, these dependencies show a mirror behavior with respect to $\theta = 30^\circ$. This means that from given FLA and FTA positions, two values are possible: $\theta \in [0,30]^\circ$ or its mirror $60^\circ - \theta$. As a consequence, θ will be given in the range of 0–30° in all plots in Figure 6 with the possibility that the values attributed to θ -2L-MoS₂ could be $60^\circ - \theta$ instead. For instance, the data from both 2Hc-2L-MoS₂ and 3R-2L-MoS₂ are reported at $\theta = 0^\circ$ in these plots. From the positions of the FTA and FLA, we claim that the spectra of the two θ -2L-MoS₂ displayed in Figure 5 correspond to 20°-2L-MoS₂ (Figure 5b–d, solid green line) and 30°-2L-MoS₂ (Figure 5a–d, solid red line), respectively.

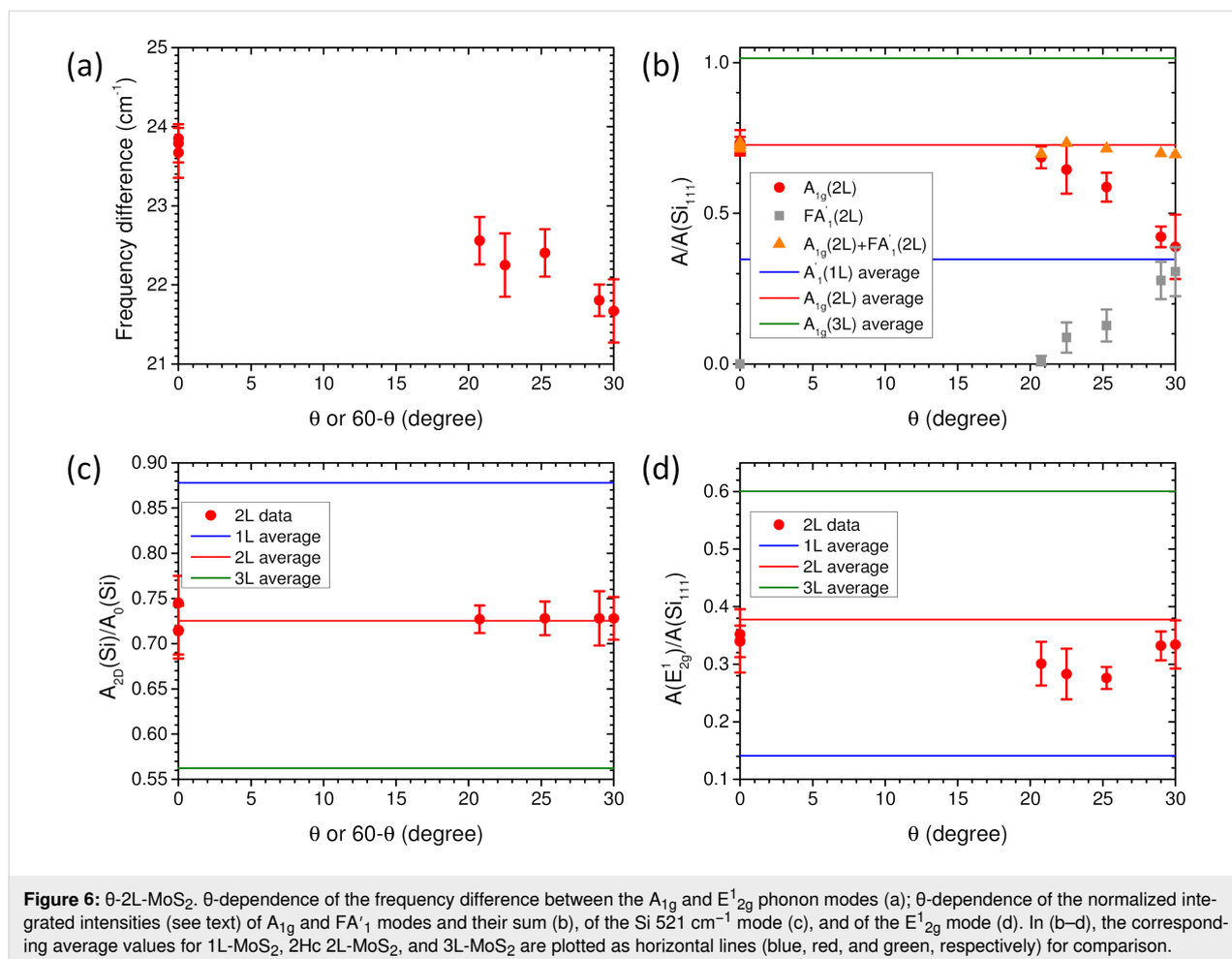
The dependence of A_{1g} and E_{2g}¹ modes on the twist angle (derived from the positions of FTA and FLA modes) is reported in Figure 5c. The frequency of the E_{2g}¹ mode in 3R-, 2Hc-, and θ -2L-MoS₂ is downshifted with respect to its frequency in 1L-MoS₂, and it does not show a clear dependence on the twist angle. In contrast, the profile of the A_{1g} mode signifi-

cantly depends on the twist angle, and a new mode, named FA'₁, appears on the high-frequency side of the A_{1g} mode. The FA'₁ mode is identified as Raman scattering from moiré phonons associated with the A'₁ dispersion curve of 1L-MoS₂. It is folded onto the zone center and, consequently, becomes Raman active [20]. Obviously, its frequency depends on the twist angle and the θ -dependence of the FA'₁ frequency was recently established both theoretically and experimentally (see Figure 3e in [20]). On the basis of these previous results, we have been able to evaluate the value of θ for each 2L-MoS₂ investigated from the position of the FA'₁ mode. The values of the angles derived from the position of FTA/FLA and FA'₁ are in close agreement.

The objective of this work is to characterize the thickness of all MoS₂ flakes. The relevance of the criteria based on the frequency ($\Delta\omega_{A-E}$) and normalized integrated intensity ($A(A_{1g})/A(Si_{111})$) of the A_{1g} mode has to be reevaluated in twisted 2L-MoS₂ flakes. As shown in Figure 5d, the normalized intensity of the 521 cm⁻¹ Si mode from the substrate underneath MoS₂ flakes is close in all the 2L-MoS₂ and independent of the twist angle.

Figure 6 summarizes and details the dependence on the twist angle of the four Raman criteria defined above for 2L-MoS₂. In all plots of Figure 6, the values of angles were derived from the positions of FTA, FLA and FA'₁. The values of the different criteria measured for θ -2L-MoS₂ are compared with the average values of the same criteria measured on exfoliated 1L-, 2L-, and 3L-MoS₂ flakes. In θ -2L-MoS₂, the value of $\Delta\omega_{A-E}$ strongly depends on the twist angle and significantly differs from the average value measured in 2Hc-2L-MoS₂ (Figure 6a). For $\theta = 30^\circ$, the value of $\Delta\omega_{A-E}$ is close to the one found in CVD 1L-MoS₂ [51]. In consequence, using $\Delta\omega_{A-E}$ alone could lead to a wrong evaluation of the thickness of twisted 2L-MoS₂.

The normalized integrated intensity $A(A_{1g})/A(Si_{111})$ significantly decreases when the twist angle increases (Figure 6b, red dots), and in 30°-2L-MoS₂, the value of $A(A_{1g})/A(Si_{111})$ is close to the average value found in 1L-MoS₂ (Figure 6b, blue solid line). The behavior of $A(A_{1g})/A(Si_{111})$ is opposite to the one of the normalized integrated intensity of the FA'₁ mode, $A(FA'₁)/A(Si_{111})$, the latter increasing with the twist angle (Figure 6b, gray squares). These results are in qualitative agreement with those reported in [40]. It can be emphasized that the integrated intensity of A_{1g} and FA'₁ bands taken together (Figure 6b, orange triangles) is close to the average value found for 2Hc-2L-MoS₂ (Figure 6b, red solid line). The reason for this compensation between $A(A_{1g})$ and $A(FA'₁)$ is not clear yet, but it could present a practical advantage in the use of the global integrated intensity of the spectral band, located around the po-



sition of the A_{1g} mode for the evaluation of the thickness of twisted MoS₂ flakes.

We also observed a tendency for $A(E_{2g}^1)/A(\text{Si}_{111})$ to be slightly lower for θ -2L-MoS₂ than for 2Hc-2L-MoS₂ (or 3R-2L-MoS₂, which is similar), but to a lesser extent compared to $A(A_{1g})/A(\text{Si}_{111})$, that is ca. 20% vs ca. 50% at maximum, respectively (Figure 6d). These results are also in qualitative agreement with those reported in [40]. Finally, only the value of the $A_{2D}(\text{Si})/A_0(\text{Si})$ ratio seems to provide a robust/reliable information to characterize the thickness of MoS₂ flakes, since it is found largely independent of θ in all measured 2L-MoS₂ samples (Figure 6c). Even if further work is needed to complete the data presented here with other values of θ and twisted MoS₂ samples with $N > 2$, we anticipate that the value of $A_{2D}(\text{Si})/A_0(\text{Si})$ ratio would be close in twisted and 2Hc-stacked MoS₂ multilayers. However, as previously recalled, the sensitivity of this ratio to the SiO₂ thickness and the gap between the $A_{2D}(\text{Si})/A_0(\text{Si})$ ratios for N and $N + 1$ permit to ensure the determination of N with a rather good reliability only for $N \leq 5$.

DLI-PP-CVD MoS₂ nanoflakes

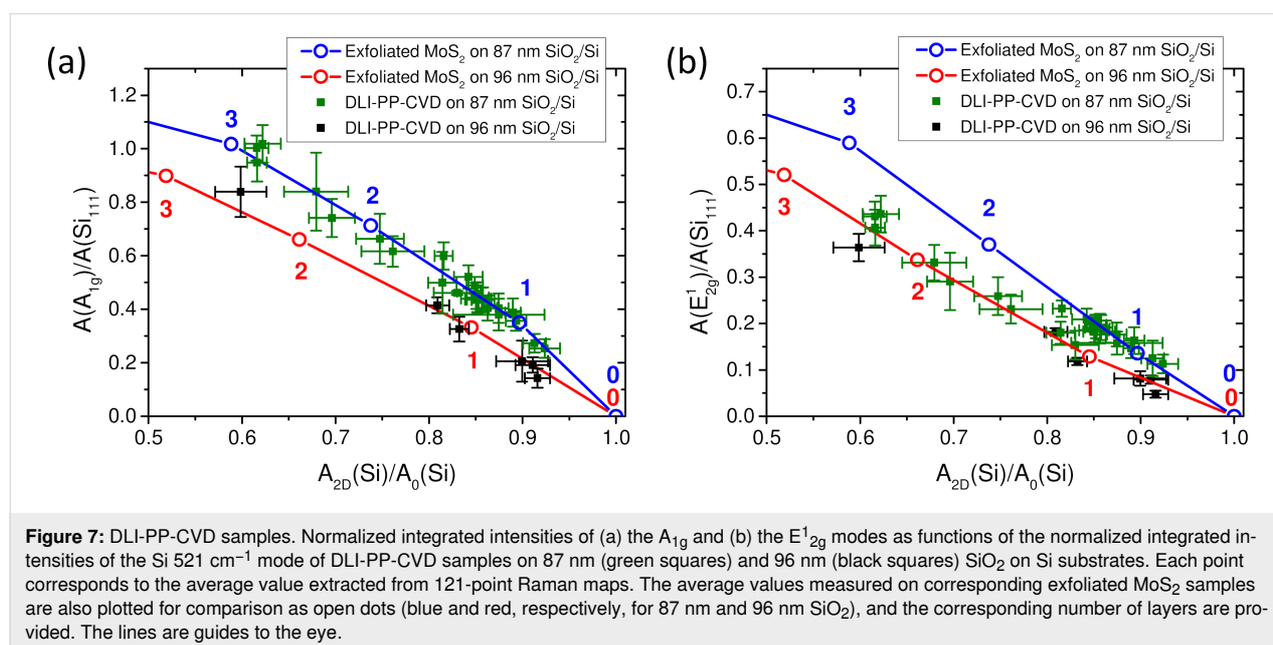
The aim of this part is to define which Raman information is relevant to estimate the thickness of MoS₂ samples produced by DLI-PP-CVD. These samples are significantly different from the previous ones (exfoliated and standard CVD). Indeed, they are constituted of nanoflakes (with a lateral size of typically 50 nm, i.e., well below the laser spot size) with possibly a distribution of thicknesses and twist angles between adjacent layers of multilayer domains and a higher number of defects (the average inter-defect distance ranges from 3 to 6 nm as estimated from the LA and A_{1g} intensity ratio [52]). In addition, the MoS₂ surface coverage is a priori unknown and can be incomplete. It is then necessary to implement a first check criterion that ensures that the thickness estimation method based on the comparison with results obtained on exfoliated samples is still valid. More generally, this point is critical for the characterization of samples synthesized using new methods or new precursors that can lead to the co-deposition of several by-products (such as carbon, oxides, and metals), which can significantly change the measured Raman intensities. Based on the results presented in the previous sections, we have shown that the value of the

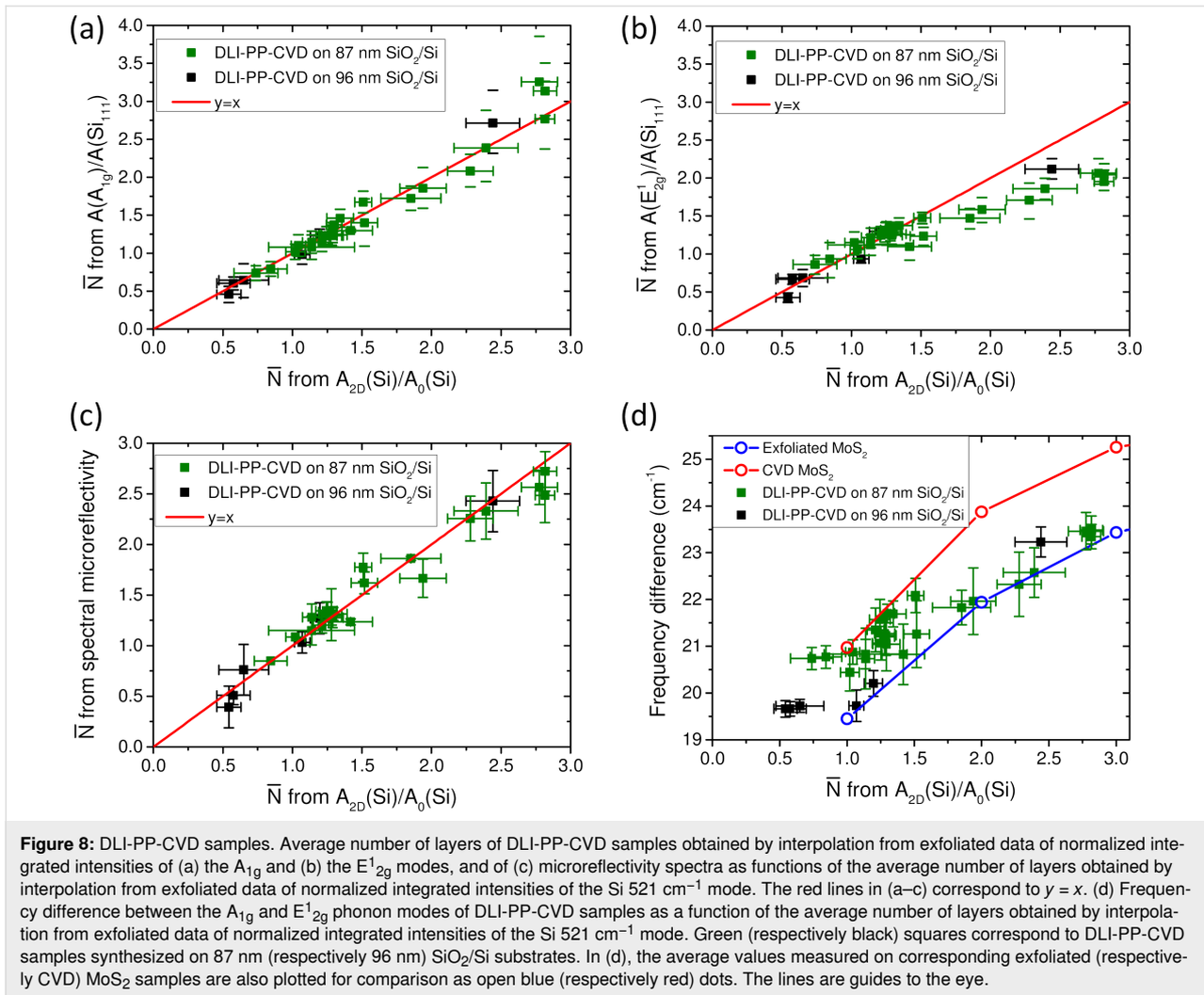
$A_{2D}(\text{Si})/A_0(\text{Si})$ ratio provides a robust/reliable Raman information to characterize the thickness of MoS_2 flakes for $N \leq 5$. However, this parameter does not rely unambiguously on the presence of MoS_2 . The deposition of any other material would influence its value and could lead to a wrong estimation. In the most general case, the sample characteristics are not perfectly known and can be significantly different from the reference characteristics. As a consequence, it seems mandatory to compare the thickness estimated from the $A_{2D}(\text{Si})/A_0(\text{Si})$ ratio with other measurands directly related to the presence of MoS_2 . To this aim, we propose to use jointly the normalized integrated intensity of the MoS_2 phonon modes, namely $A(A_{1g})/A(\text{Si}_{111})$ and/or $A(E^1_{2g})/A(\text{Si}_{111})$.

In Figure 7, the values of $A(A_{1g})/A(\text{Si}_{111})$ (Figure 7a) and $A(E^1_{2g})/A(\text{Si}_{111})$ (Figure 7b) are plotted as functions of the value of $A_{2D}(\text{Si})/A_0(\text{Si})$. In these graphs, the data obtained on DLI-PP-CVD samples are compared with the average reference measurements established previously on exfoliated MoS_2 deposited on Si/SiO_2 substrate with the same SiO_2 thicknesses, namely 96 nm (red open dots in Figure 7) and 87 nm (blue open dots in Figure 7). Note that in the exfoliated samples, the exact number of layers N is perfectly known and given on the plots of Figure 7 close to corresponding open dots. The idea behind this representation comes from the expectation that the presence of contaminations or deposition of others species would have a different impact on the Raman intensity coming from MoS_2 in the film and on the one coming from the Si substrate underneath the deposited thin film. It is, thus, expected that the measurements on contaminated or highly defective MoS_2 thin films will fall off the reference curve. Indeed, data obtained on poorly

crystalline MoS_2 films synthesized by DLI atomic layer deposition (not shown) are found systematically and significantly below the corresponding reference curve. Concerning the DLI-PP-CVD samples presented in Figure 7, the $A(A_{1g})/A(\text{Si}_{111})$ vs $A_{2D}(\text{Si})/A_0(\text{Si})$ dependence is found fully compatible with the respective reference exfoliated curves (Figure 7a). The $A(E^1_{2g})/A(\text{Si}_{111})$ vs $A_{2D}(\text{Si})/A_0(\text{Si})$ data points mainly agree for thin layers ($A_{2D}(\text{Si})/A_0(\text{Si}) > 0.8$) but fall systematically below the corresponding reference exfoliated curves for thicker layers ($0.8 > A_{2D}(\text{Si})/A_0(\text{Si}) > 0.6$) as shown in Figure 7b.

Another way to compare the results is estimating the thickness of DLI-PP-CVD samples by interpolation from exfoliated data of the measured values for $A_{2D}(\text{Si})/A_0(\text{Si})$, $A(A_{1g})/A(\text{Si}_{111})$, and $A(E^1_{2g})/A(\text{Si}_{111})$. In Figure 8a (respectively 8b), the average number of layers (\bar{N}) obtained using $A(A_{1g})/A(\text{Si}_{111})$ (respectively $A(E^1_{2g})/A(\text{Si}_{111})$) are plotted as a function of the number derived from $A_{2D}(\text{Si})/A_0(\text{Si})$. It can be emphasized that non-integer values are found for \bar{N} , indicating the presence of a mix with unknown proportions of bare substrate (0L), 1L- MoS_2 , 2L- MoS_2 , 3L- MoS_2 , and so on in the investigated DLI-PP-CVD films. It is also noticeable that the errors of \bar{N} estimated from $A(A_{1g})/A(\text{Si}_{111})$ become larger when \bar{N} is close to 3 as a consequence of the smoother dependence of this parameter with \bar{N} . In agreement with the conclusion drawn above from Figure 7, Figure 8a illustrates the coherence between the values of \bar{N} derived from $A(A_{1g})/A(\text{Si}_{111})$ and $A_{2D}(\text{Si})/A_0(\text{Si})$. All data remain close to the red solid line that represents the ideal relation $y[\bar{N} \text{ via } A(A_{1g})/A(\text{Si}_{111})] = x[\bar{N} \text{ via } A_{2D}(\text{Si})/A_0(\text{Si})]$. Figure 8b as well confirms that the values of \bar{N} derived from $A(E^1_{2g})/A(\text{Si}_{111})$ and $A_{2D}(\text{Si})/A_0(\text{Si})$ agree well for $\bar{N} < 1.5$, but





the values of \bar{N} from $A(E_{2g}^1)/A(\text{Si}_{111})$ are systematically lower than those obtained from $A_{2D}(\text{Si})/A_0(\text{Si})$ when $\bar{N} > 1.5$. One explanation could be the presence of a larger proportion of multi-layer regions in the thicker samples, for which, if they are twisted, $A(E_{2g}^1)/A(\text{Si}_{111})$ has been shown to be attenuated in the previous section. If so, the question then arises why the same behavior is not observed for $A(A_{1g})/A(\text{Si}_{111})$ contrary to what would be expected. A possibility could be that because of the observed broadening of the A_{1g} mode in DLI-PP-CVD samples (presumably due to local heterogeneities in terms of doping, strain, defects, or thickness), the FA'_{1g} mode becomes indistinguishable from the A_{1g} mode. As a consequence, the intensity of FA'_{1g} would merge with $A(A_{1g})$ and compensate its attenuation. Other explanations relying on the presence of defects or strain cannot be disregarded, and further works are needed to fully clarify this point.

In order to further confirm the validity of the estimations of \bar{N} for DLI-PP-CVD samples, we compare in Figure 8c the \bar{N}

values derived from $A_{2D}(\text{Si})/A_0(\text{Si})$ with the ones obtained independently from spectral microreflectivity. A very good agreement is found between the two series of data, establishing definitively the relevance of the $A_{2D}(\text{Si})/A_0(\text{Si})$ ratio to give with good accuracy the average thickness of DLI-PP-CVD MoS_2 samples for $\bar{N} \leq 3$. This agreement justifies the use of the values of \bar{N} derived from $A_{2D}(\text{Si})/A_0(\text{Si})$ as abscissa axis in the previous plots.

Finally, in Figure 8d the frequency difference between the A_{1g} and E_{2g}^1 phonons is plotted as a function of \bar{N} estimated from $A_{2D}(\text{Si})/A_0(\text{Si})$ for DLI-PP-CVD samples and compared to the data obtained on exfoliated and CVD MoS_2 . DLI-PP-CVD data are distributed between the two curves obtained from the reference samples. This further confirms that this measurand cannot be used to evaluate with good accuracy their average thicknesses. Nevertheless this comparison can be informative, showing that samples with $\bar{N} < 1$ are most certainly mainly composed of 1L- MoS_2 and suggesting that the proportions of

2L-MoS₂, 3L-MoS₂, or more gradually increase with \bar{N} , which is compatible with AFM observations (not shown).

To get further insight on the number of layer distributions in DLI-PP-CVD samples, we have measured their ULF modes. Representative ULF spectra are shown in Figure 9a for samples with average thicknesses ranging from 0.6 up to 2.8 MoS₂ layers as estimated from $A_{2D}(\text{Si})/A_0(\text{Si})$. Up to $\bar{N} = 1.3$, only the LB mode of 2L-MoS₂ is observed around 40 cm⁻¹ [20,38–41], showing that these samples can only be composed of 1L-MoS₂ and twisted 2L-MoS₂ plus possibly uncovered (bare substrate) regions. For thicker samples, the S mode of 2L-MoS₂ around 24 cm⁻¹ is additionally visible, as well as a signal between 25 and 30 cm⁻¹, corresponding to the LB and S modes of 3L-MoS₂ [32,33]. For $\bar{N} \geq 2.4$, the LB mode of 4L-MoS₂ is also present around 21 cm⁻¹; there may also be a weak signal around 17 cm⁻¹ (corresponding to the LB mode of 5L-MoS₂) reflecting the presence of 5L-MoS₂. The S mode of 4L-MoS₂ could be present as well around 28 cm⁻¹, but it is hardly distinguishable from the LB and S modes of 3L-MoS₂. Thus, ULF Raman spectra give valuable qualitative information on the different N present in each sample. Quantitative information relies on the determination of the surface coverages for each N (σ_N), that is, the ratio between the surface covered by exactly N layers and the total surface. With $N = 0$ standing for the bare substrate and N_{max} being the largest number of layers present in the sample, the definition of the average number of layers \bar{N} can be written as

$$\bar{N} = \sum_{N=1}^{N_{\text{max}}} \sigma_N \times N \quad (1)$$

and the total coverage (including bare substrate areas) is obviously 100%:

$$\sum_{N=0}^{N_{\text{max}}} \sigma_N = 1 \quad (2)$$

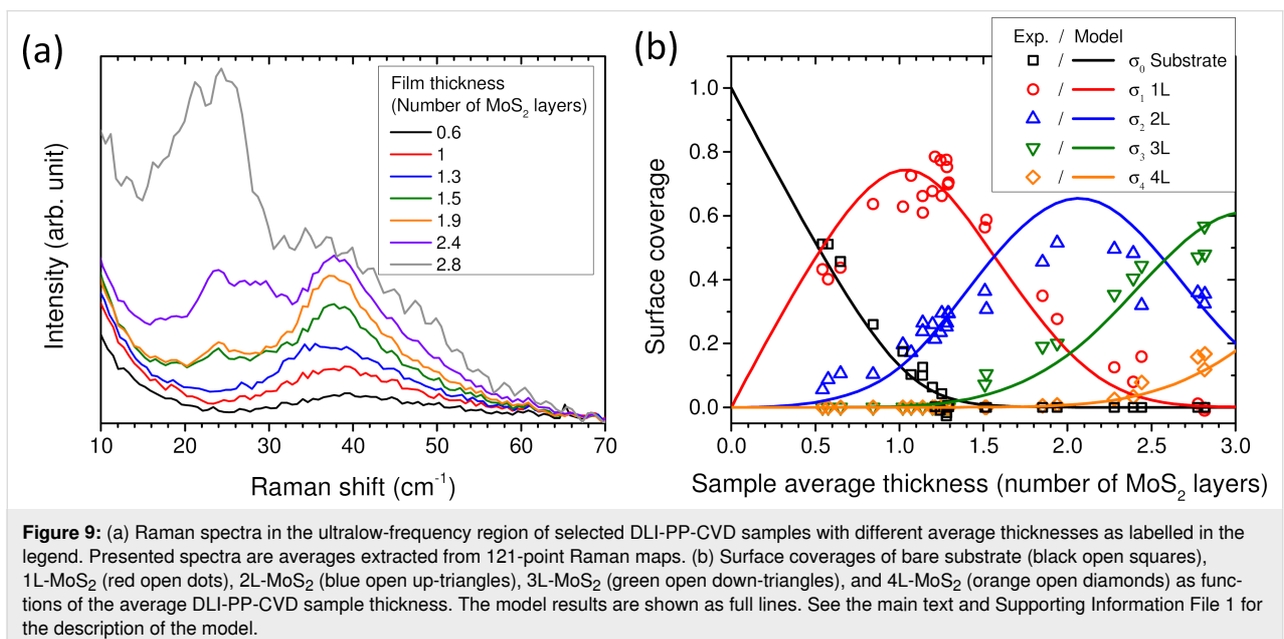
AFM imaging (see Supporting Information File 1, Figure S4) reveals that for $\bar{N} > 1.25$, the surface is fully covered by MoS₂, that is, $\sigma_0 = 0$, which removes an unknown. In addition, for $\bar{N} < 1.3$, there is no signature of more than two layers, and we can set $\sigma_{N \geq 3} = 0$ with confidence. Hence, for $1.25 < \bar{N} < 1.3$, the set of Equations 1 and 2 simplifies to

$$\begin{aligned} \bar{N} &= \sigma_1 \times 1 + \sigma_2 \times 2 \\ \sigma_1 + \sigma_2 &= 1 \end{aligned}$$

This allows one to readily determine the two remaining unknowns σ_1 and σ_2 , since \bar{N} is known from $A_{2D}(\text{Si})/A_0(\text{Si})$.

Hereafter, a linear relationship between the Raman signal $I_{\text{LB}}^{2\text{L}}$, which is the LB mode peak intensity of 2L ($N = 2$) areas (the broad but well-identified 40 cm⁻¹ peak), and the surface coverage is assumed, namely $\sigma_2 = \alpha_2 I_{\text{LB}}^{2\text{L}}$. The ratio $\alpha_2 = \sigma_2 / I_{\text{LB}}^{2\text{L}}$ is determined from five samples ($1.25 < \bar{N} < 1.3$) for which we now have both the coverage σ_2 and the Raman signal $I_{\text{LB}}^{2\text{L}}$.

Because α_2 is now known and assuming that the linearity between σ_2 and the Raman signal $I_{\text{LB}}^{2\text{L}}$ holds (which should be a good approximation for the thin multilayers considered here), $\sigma_2 = \alpha_2 I_{\text{LB}}^{2\text{L}}$ can be obtained directly for all samples from the Raman spectra, and is thus no longer an unknown.



We now turn to the samples with $\bar{N} < 1.25$, which may present some bare substrate areas, so σ_0 and σ_1 are a priori unknown. Both \bar{N} and σ_2 are determined as explained above, and $\sigma_{N \geq 3} = 0$ is again a safe estimate. Hence, Equations 1 and 2 reduce to the system

$$\begin{aligned}\bar{N} &= \sigma_1 \times 1 + \sigma_2 \times 2 \\ \sigma_0 + \sigma_1 + \sigma_2 &= 1\end{aligned}$$

which may be solved trivially for σ_0 and σ_1 .

A similar approach can be used for samples with $1.3 < \bar{N} < 2$ that are fully covered ($\sigma_0 = 0$) and might present some trilayers but show no trace of thicker layers. We set $\sigma_{N \geq 4} = 0$, and the system of equations reduces to

$$\begin{aligned}\bar{N} &= \sigma_1 \times 1 + \sigma_2 \times 2 + \sigma_3 \times 3 \\ \sigma_1 + \sigma_2 + \sigma_3 &= 1\end{aligned}$$

Both \bar{N} and σ_2 are determined as explained above. Thus, the system can be trivially solved for the two remaining unknowns σ_1 and σ_3 .

At this point it would be natural to get the proportionality between σ_3 and a Raman signal attributed to 3L areas ($N = 3$), and proceed recursively to obtain σ_4 in slightly thicker layers, and so on and so forth. In practice this becomes challenging because of the uncertainty on the 3L ($N = 3$) Raman signal, which is less clear than the 2L ($N = 2$) peak. Another approach gave better results.

Three samples with \bar{N} between 2.75 and 2.85 are thick enough to neglect σ_0 and σ_1 , yet thin enough for σ_5 to also be negligible as a first approximation. Equations 1 and 2 then reduce to

$$\begin{aligned}\bar{N} &= \sigma_2 \times 2 + \sigma_3 \times 3 + \sigma_4 \times 4 \\ \sigma_2 + \sigma_3 + \sigma_4 &= 1\end{aligned}$$

where \bar{N} and σ_2 are known, so σ_3 and σ_4 can be determined readily.

The LB mode of 4L-MoS₂, located around 21 cm⁻¹, is sufficiently separated from other modes to be identified (which was not the case for I_{LB}^3), so that I_{LB}^4 can be extracted from the spectra. From the three $2.75 < \bar{N} < 2.85$ samples $\alpha_4 = \sigma_4 / I_{\text{LB}}^4$ is determined. Then, assuming again a linear relationship $\sigma_4 = \alpha_4 I_{\text{LB}}^4$, the coverage by 4L ($N = 4$) layers can be determined for all samples. This removes another unknown.

Now the last remaining case of $2 < \bar{N} < 2.75$ samples can be solved, as

$$\begin{aligned}\bar{N} &= \sigma_1 \times 1 + \sigma_2 \times 2 + \sigma_3 \times 3 + \sigma_4 \times 4 \\ \sigma_1 + \sigma_2 + \sigma_3 + \sigma_4 &= 1\end{aligned}$$

give σ_1 and σ_3 directly, since \bar{N} , σ_2 and σ_4 are known.

The results obtained using this procedure are shown in Figure 9b where σ_N (with N from 0 to 4) is plotted as a function of \bar{N} , the average sample thickness. On this graph, all values of σ_2 (respectively σ_4) are calculated using $I_{\text{LB}}^{2\text{L}}$ (respectively $I_{\text{LB}}^{4\text{L}}$) even for the samples used to derive the proportionality coefficient α_2 (respectively α_4). As shown in Figure 9b for samples with $1.25 < \bar{N} < 1.3$ (respectively $2.75 < \bar{N} < 2.85$), we find by this way $-0.03 < \sigma_0 < 0.04$ (respectively $-0.01 < \sigma_1 < 0.01$) with little fluctuations around the expected value of 0.

Just below the full coverage of the sample surface by MoS₂ ($\sigma_0 > 0$), both σ_1 and σ_2 increase with a slight tendency of σ_2 to increase faster than σ_1 . Indeed, 1L-MoS₂ represents 80–90% of the deposited MoS₂ for $\bar{N} = 0.5$ and 70–80% for $\bar{N} = 1.3$. The maximum of σ_1 is reached around $\bar{N} = 1.3$ when the sample surface is totally covered by MoS₂ ($\sigma_0 = 0$), and σ_1 starts to decrease above this value. Around $\bar{N} = 1.6$, 1L-MoS₂ only represents 50% of the MoS₂. σ_2 continues to increase and reaches a maximum value of $\approx 50\%$ around $\bar{N} = 2$ and then decreases for thicker samples. 3L-MoS₂ starts to appear after the substrate surface is completely covered by MoS₂ and increases continuously, representing about 50% of the thickest samples ($\bar{N} \approx 2.8$).

In order to verify our approach, we implemented a 2D growth toy model (see Supporting Information File 1 for details). The model results are shown in Figure 9b as full lines and give a good agreement with the experimental results. It should be noted that within this representation ($\sigma_N = f(\bar{N})$), the results of the model are remarkably robust to any parameter changes (the curves are almost insensitive to either doubling or halving the cell size and, thus, the advance rate, or to multiplying or dividing the growth rate by 5). In other words, this means that this comparison with the experiment cannot be used to validate any model parameters but demonstrates the relevance of the proposed procedure to estimate the σ_N from the experiments. Nevertheless, it has to be noted that while for $\bar{N} < 1.3$ the ULF Raman signature of 2L-MoS₂ remains very similar, it is not the case for thicker samples with the notable appearance of the S mode of 2L-MoS₂ around 24 cm⁻¹ [32,33]. This could mean that the stacking order distribution changes. As a consequence, the hypothesis based on the proportionality between $I_{\text{LB}}^{2\text{L}}$ and σ_2 would probably be less valid above $\bar{N} = 1.3$, and an error on the absolute values deduced can be expected. However, the appearance of the S mode of 2L-MoS₂ around 24 cm⁻¹ could also be related to $N\text{L-MoS}_2$ (with $N \geq 3$) constituted of a stacking se-

quence where 2L are not twisted, for example, the so-called $t(1+2)L$, $t(2+2)L$, ... structures [26,53]. In this case, our hypothesis would remain more appropriate. Despite this unknown as well as the other approximations made, we believe that the main tendencies can still be captured by the proposed analysis. Further works are needed to determine and improve the accuracy of the method.

Conclusion

In this work we have reviewed all Raman information leading to the evaluation of the thickness of MoS₂ flakes, that is, the layer number N . First, we have analyzed in detail the effects of some experimental parameters, namely the wavelength of the incident laser light used in the experiments, the power of the incident light, and the oxide thickness of the SiO₂/Si substrate on which the flakes are deposited, on the quality and accuracy of Raman results. Based on this analysis, an experimental protocol has been defined and systematically applied to large MoS₂ flakes (i.e., single-domain flakes much larger than the laser spot), including twisted MoS₂ flakes, prepared by different methods on the one hand and to MoS₂ thin films composed of nanoflakes prepared by the DLI-PP-CVD method on the other hand. Special attention was paid to the measurement statistics.

The limits of different Raman criteria which allow one to determine the thicknesses of MoS₂ flakes, namely (i) the value of $\Delta\omega_{A-E}$, (ii) the value of the normalized integrated intensity of A_{1g} and E_{2g}^1 MoS₂ modes, and (iii) the value of the $A_{2D}(Si)/A_0(Si)$ ratio, have been precisely studied in the different types of MoS₂ samples. We definitely confirm that $\Delta\omega_{A-E}$ cannot be considered a robust criterion to derive the number of layers in MoS₂ samples. We found that the value of the $A_{2D}(Si)/A_0(Si)$ ratio provides the most robust/reliable information to characterize the thickness of MoS₂ large flakes, especially since it is found largely independent of the twist angle. The limit of application of this criterion is $N \leq 5$, under the condition that the SiO₂ thickness is precisely known.

We then apply this analysis procedure to DLI-PP-CVD samples constituted of nanoflakes with a lateral size of typically 50 nm (well below the laser spot size) with possibly a distribution of thicknesses and twist angles between adjacent layers of multilayer domains and a higher number of defects. Our results definitively establish the relevance of the $A_{2D}(Si)/A_0(Si)$ ratio to give with good accuracy their average thickness \bar{N} , for $\bar{N} \leq 3$. Nevertheless, we emphasize that this criterion is not only related to the presence of MoS₂ and can be influenced by several factors, such as the co-deposition of by-products or the presence of defects, leading to a wrong estimation of \bar{N} . We propose to combine $A_{2D}(Si)/A_0(Si)$ with the normalized inte-

grated intensity of the MoS₂ phonon modes, namely $A(A_{1g})$ and/or $A(E_{2g}^1)$. Although limiting the application to $\bar{N} \leq 3$, this approach enables the validation of the $A_{2D}(Si)/A_0(Si)$ ratio to determine \bar{N} in the presented case, and we anticipate that it would avoid possible errors in unfavorable situations.

Finally, to get further insight on the number of layer distributions in DLI-PP-CVD samples, we have measured their ULF modes. An original procedure based of the measurement of the intensity of the layer breathing modes allows one to evaluate the surface coverage (σ_N) for each N . A 2D growth toy model gives a good agreement with the experimental results supporting the proposed procedure to estimate the σ_N from the ULF spectra.

Experimental

Samples preparation

Mechanical exfoliation

MoS₂ flakes were obtained by micromechanical cleavage of a MoS₂ crystal (HQ graphene) using scotch tape (Nitto) and PDMS slabs (Gel-pak). They were then transferred onto Si substrates with SiO₂ layers of different thicknesses, namely 84, 87, 90, and 96 nm. Flakes were selected by optical microscopy and their thicknesses were determined by optical contrast.

Standard CVD process

MoS₂ was grown by CVD on 87 nm SiO₂ on Si substrates using MoO₃ (Sigma-Aldrich, 25 mg) and sulfur (Sigma-Aldrich, 250 mg) powders as solid precursors using a 1 inch quartz tube furnace. MoO₃ powder was placed in the center of the heating zone of the furnace, while sulfur was placed upstream at the furnace inlet. Prior to growth, air was evacuated by flowing Ar (ultrahigh purity, Linde) for 15 min at 200 sccm, after which the tube was heated to 200 °C for 10 min. The temperature was then increased to 750 °C under Ar (100 sccm), and it was held at this value for 15 min before cooling naturally to room temperature.

Direct-liquid injection pulsed-pressure chemical vapor deposition (DLI-PP-CVD)

The 12 × 11 mm SiO₂/Si (with 87 nm or 96 nm SiO₂ thicknesses) substrates were cleaned in acetone (C₃H₆O, technical, Acros Organics), isopropanol (C₃H₇OH, 99.8%, Höfer Chemie GmbH) and deionized water (H₂O, Acros Organics) under ultrasonic agitation for 10 min each, before being blown dry with nitrogen. They were then immediately loaded on the susceptor of the reaction chamber (Annealsys MC-050) for deposition. Solutions of 0.001 M molybdenum hexacarbonyl (Mo(CO)₆, 98%, Strem Chemicals) and 0.002 M sulfur (S, 99.999%, Acros Organics) in anhydrous toluene (C₆H₅CH₃, 99.8%, Sigma-Aldrich) were used as precursors. The process is

as follows: Following sample installation, the chamber is closed and brought to about 0.02 mbar. For monolayer depositions, it is imperative that the substrate is thoroughly cleaned and free of adsorbates. Therefore, to ensure complete desorption of remaining contaminants, the samples were kept for 30 min under vacuum at room temperature inside the deposition chamber. For the first part of the process, the pumping direction is reversed so that all species are pumped from the deposition chamber to the back of the reactor.

Nitrogen (800 sccm) is flowed through the chamber (200 sccm through the gas line, and 600 sccm through two injection heads) and the substrates are brought to 750 °C at a ramp of 2 °C/s. The reactor is kept in this state for 5 min for homogenization purposes. While still in reverse direction pumping, 0.3 g/min of both precursors are injected and vaporized to prepare the evaporation system for deposition. Then, the Mo(CO)₆ injection is stopped, the pumping direction is switched back to the deposition direction and hydrogen (40 sccm) is added to the gas mix. For 1 min, sulfur is injected to clean any remaining contaminants, and to prepare the surface of the substrate for MoS₂ deposition, then the deposition works in 20 s cycles. During one cycle, a single pulse of 3 to 10 ms of Mo(CO)₆ is injected while the S injection is set to 0.3 g/min. This 20 s cycle is repeated 80 to 160 times. The quantity of MoS₂ deposited is controlled by the quantity of Mo(CO)₆ injected, that is, the pulse duration and the number of cycles.

Raman spectroscopy

Raman spectra and maps were recorded using an Acton spectrometer fitted with a Pylon CCD detector and a 1800 grooves/mm grating ($\approx 0.6 \text{ cm}^{-1}$ between each CCD pixel). The samples were excited with a 532 nm (2.33 eV) laser (Newport Millennia Prime or Cobolt Samba) through an Olympus microscope objective either 100× (numerical aperture 0.9) or 50× (numerical aperture 0.5). The full width at half-maximum (FWHM) of the focused laser spot with the 100× objective is about 380 nm. Optimized focus conditions were checked for each measurement. The samples were mounted on a three-axis piezoelectric stage (Physik Instrumente) to ensure the precise positioning and focusing of the laser spot. A Si(111) wafer with only native oxide sample was used as a daily reference for the system. The laser power was continuously measured during acquisitions allowing for intensity normalizations of the Raman spectra at each point of the maps. All data presented in this paper, unless specified otherwise, are extracted from Raman maps constituted by hundreds to thousands points (see Supporting Information File 1 for an example), which were analyzed using a custom-made software. All reported points are the average values obtained by Gaussian fitting of the data distribution extracted from Raman maps (corresponding to

hundreds to several thousands of spectra), and the error bars correspond to 99.7% confidence intervals (± 3 standard deviations).

2D growth toy model

The model used the DynamicGrids.jl package, which was part of the Dispersal.jl framework [54], see Supporting Information File 1 for more details.

Supporting Information

Supporting Information File 1 contains additional figures with an example of Raman maps, the Si mode as a function of the laser power, a comparison between two microscope objectives, other intensity references, atomic force microscopy images, and details of the 2D growth toy model. Supporting Information File 2 is a recording of the growth simulation.

Supporting Information File 1

Additional experimental data.

[<https://www.beilstein-journals.org/bjnano/content/supplementary/2190-4286-15-26-S1.pdf>]

Supporting Information File 2

Recording of the growth simulation.

[<https://www.beilstein-journals.org/bjnano/content/supplementary/2190-4286-15-26-S2.mp4>]

Funding

This work was supported by Occitania region “Recherche et Société(s)” program in the frame of DIMENSION project N°2018-003267.

ORCID® iDs

Vincent Astié - <https://orcid.org/0000-0001-6548-178X>

Romain Parret - <https://orcid.org/0000-0003-0662-6373>

Sylvie Contreras - <https://orcid.org/0000-0002-4733-9998>

Matthieu Paillet - <https://orcid.org/0000-0001-7038-9101>

References

- Novoselov, K. S.; Jiang, D.; Schedin, F.; Booth, T. J.; Khotkevich, V. V.; Morozov, S. V.; Geim, A. K. *Proc. Natl. Acad. Sci. U. S. A.* **2005**, *102*, 10451–10453. doi:10.1073/pnas.0502848102
- Gibney, E. *Nature* **2015**, *522*, 274–276. doi:10.1038/522274a
- Lin, Z.; McCreary, A.; Briggs, N.; Subramanian, S.; Zhang, K.; Sun, Y.; Li, X.; Borys, N. J.; Yuan, H.; Fullerton-Shirey, S. K.; Chernikov, A.; Zhao, H.; McDonnell, S.; Lindenberg, A. M.; Xiao, K.; LeRoy, B. J.; Drndić, M.; Hwang, J. C. M.; Park, J.; Chhowalla, M.; Schaak, R. E.; Javey, A.; Hersam, M. C.; Robinson, J.; Terrones, M. *2D Mater.* **2016**, *3*, 042001. doi:10.1088/2053-1583/3/4/042001

4. Withers, F.; Del Pozo-Zamudio, O.; Mishchenko, A.; Rooney, A. P.; Gholinia, A.; Watanabe, K.; Taniguchi, T.; Haigh, S. J.; Geim, A. K.; Tartakovskii, A. I.; Novoselov, K. S. *Nat. Mater.* **2015**, *14*, 301–306. doi:10.1038/nmat4205
5. Jariwala, D.; Sangwan, V. K.; Lauhon, L. J.; Marks, T. J.; Hersam, M. C. *ACS Nano* **2014**, *8*, 1102–1120. doi:10.1021/nn500064s
6. Georgiou, T.; Jalil, R.; Belle, B. D.; Britnell, L.; Gorbachev, R. V.; Morozov, S. V.; Kim, Y.-J.; Gholinia, A.; Haigh, S. J.; Makarovskiy, O.; Eaves, L.; Ponomarenko, L. A.; Geim, A. K.; Novoselov, K. S.; Mishchenko, A. *Nat. Nanotechnol.* **2013**, *8*, 100–103. doi:10.1038/nnano.2012.224
7. Wang, H.; Yu, L.; Lee, Y.-H.; Shi, Y.; Hsu, A.; Chin, M. L.; Li, L.-J.; Dubey, M.; Kong, J.; Palacios, T. *Nano Lett.* **2012**, *12*, 4674–4680. doi:10.1021/nl302015v
8. Baugher, B. W. H.; Churchill, H. O. H.; Yang, Y.; Jarillo-Herrero, P. *Nat. Nanotechnol.* **2014**, *9*, 262–267. doi:10.1038/nnano.2014.25
9. Zhang, Y. J.; Oka, T.; Suzuki, R.; Ye, J. T.; Iwasa, Y. *Science* **2014**, *344*, 725–728. doi:10.1126/science.1251329
10. Wang, Q. H.; Kalantar-Zadeh, K.; Kis, A.; Coleman, J. N.; Strano, M. S. *Nat. Nanotechnol.* **2012**, *7*, 699–712. doi:10.1038/nnano.2012.193
11. Calman, E. V.; Fogler, M. M.; Butov, L. V.; Hu, S.; Mishchenko, A.; Geim, A. K. *Nat. Commun.* **2018**, *9*, 1895. doi:10.1038/s41467-018-04293-7
12. Rivera, P.; Schaibley, J. R.; Jones, A. M.; Ross, J. S.; Wu, S.; Aivazian, G.; Klement, P.; Seyler, K.; Clark, G.; Ghimire, N. J.; Yan, J.; Mandrus, D. G.; Yao, W.; Xu, X. *Nat. Commun.* **2015**, *6*, 6242. doi:10.1038/ncomms7242
13. Mak, K. F.; Lee, C.; Hone, J.; Shan, J.; Heinz, T. F. *Phys. Rev. Lett.* **2010**, *105*, 136805. doi:10.1103/physrevlett.105.136805
14. Splendiani, A.; Sun, L.; Zhang, Y.; Li, T.; Kim, J.; Chim, C.-Y.; Galli, G.; Wang, F. *Nano Lett.* **2010**, *10*, 1271–1275. doi:10.1021/nl903868w
15. Huang, S.; Ling, X.; Liang, L.; Kong, J.; Terrones, H.; Meunier, V.; Dresselhaus, M. S. *Nano Lett.* **2014**, *14*, 5500–5508. doi:10.1021/nl5014597
16. Scheuschner, N.; Ochedowski, O.; Kaulitz, A.-M.; Gillen, R.; Schleberger, M.; Maultzsch, J. *Phys. Rev. B* **2014**, *89*, 125406. doi:10.1103/physrevb.89.125406
17. Eda, G.; Yamaguchi, H.; Voiry, D.; Fujita, T.; Chen, M.; Chhowalla, M. *Nano Lett.* **2011**, *11*, 5111–5116. doi:10.1021/nl201874w
18. Ribeiro-Soares, J.; Almeida, R. M.; Barros, E. B.; Araujo, P. T.; Dresselhaus, M. S.; Caçado, L. G.; Jorio, A. *Phys. Rev. B* **2014**, *90*, 115438. doi:10.1103/physrevb.90.115438
19. Wilson, J. A.; Yoffe, A. D. *Adv. Phys.* **1969**, *18*, 193–335. doi:10.1080/00018736900101307
20. Lin, M.-L.; Tan, Q.-H.; Wu, J.-B.; Chen, X.-S.; Wang, J.-H.; Pan, Y.-H.; Zhang, X.; Cong, X.; Zhang, J.; Ji, W.; Hu, P.-A.; Liu, K.-H.; Tan, P.-H. *ACS Nano* **2018**, *12*, 8770–8780. doi:10.1021/acsnano.8b05006
21. Cao, Y.; Fatemi, V.; Fang, S.; Watanabe, K.; Taniguchi, T.; Kaxiras, E.; Jarillo-Herrero, P. *Nature* **2018**, *556*, 43–50. doi:10.1038/nature26160
22. Yankowitz, M.; Chen, S.; Polshyn, H.; Zhang, Y.; Watanabe, K.; Taniguchi, T.; Graf, D.; Young, A. F.; Dean, C. R. *Science* **2019**, *363*, 1059–1064. doi:10.1126/science.aav1910
23. Autere, A.; Jussila, H.; Dai, Y.; Wang, Y.; Lipsanen, H.; Sun, Z. *Adv. Mater. (Weinheim, Ger.)* **2018**, *30*, 1705963. doi:10.1002/adma.201705963
24. Hsu, W.-T.; Zhao, Z.-A.; Li, L.-J.; Chen, C.-H.; Chiu, M.-H.; Chang, P.-S.; Chou, Y.-C.; Chang, W.-H. *ACS Nano* **2014**, *8*, 2951–2958. doi:10.1021/nn500228r
25. Yu, H.; Liu, G.-B.; Tang, J.; Xu, X.; Yao, W. *Sci. Adv.* **2017**, *3*, e1701696. doi:10.1126/sciadv.1701696
26. Zhang, X.; Qiao, X.-F.; Shi, W.; Wu, J.-B.; Jiang, D.-S.; Tan, P.-H. *Chem. Soc. Rev.* **2015**, *44*, 2757–2785. doi:10.1039/c4cs00282b
27. Astié, V.; Wasem-Klein, F.; Makhoul, H.; Paillet, M.; Huntzinger, J.-R.; Sauvajol, J.-L.; Zahab, A.-A.; Juillaguet, S.; Contreras, S.; Voiry, D.; Landois, P., in press.
28. Li, S.-L.; Miyazaki, H.; Song, H.; Kuramochi, H.; Nakaharai, S.; Tsukagoshi, K. *ACS Nano* **2012**, *6*, 7381–7388. doi:10.1021/nn3025173
29. Lee, C.; Yan, H.; Brus, L. E.; Heinz, T. F.; Hone, J.; Ryu, S. *ACS Nano* **2010**, *4*, 2695–2700. doi:10.1021/nn1003937
30. Molina-Sanchez, A.; Wirtz, L. *Phys. Rev. B* **2011**, *84*, 155413. doi:10.1103/physrevb.84.155413
31. Li, X.-L.; Qiao, X.-F.; Han, W.-P.; Zhang, X.; Tan, Q.-H.; Chen, T.; Tan, P.-H. *Nanotechnology* **2016**, *27*, 145704. doi:10.1088/0957-4484/27/14/145704
32. Zhao, Y.; Luo, X.; Li, H.; Zhang, J.; Araujo, P. T.; Gan, C. K.; Wu, J.; Zhang, H.; Quek, S. Y.; Dresselhaus, M. S.; Xiong, Q. *Nano Lett.* **2013**, *13*, 1007–1015. doi:10.1021/nl304169w
33. Zhang, X.; Han, W. P.; Wu, J. B.; Milana, S.; Lu, Y.; Li, Q. Q.; Ferrari, A. C.; Tan, P. H. *Phys. Rev. B* **2013**, *87*, 115413. doi:10.1103/physrevb.87.115413
34. Lee, J.-U.; Cheong, H. J. *Raman Spectrosc.* **2018**, *49*, 66–75. doi:10.1002/jrs.5200
35. Drapcho, S. G.; Kim, J.; Hong, X.; Jin, C.; Shi, S.; Tongay, S.; Wu, J.; Wang, F. *Phys. Rev. B* **2017**, *95*, 165417. doi:10.1103/physrevb.95.165417
36. Carvalho, B. R.; Malard, L. M.; Alves, J. M.; Fantini, C.; Pimenta, M. A. *Phys. Rev. Lett.* **2016**, *116*, 089904. doi:10.1103/physrevlett.116.089904
37. Nemanich, R. J.; Tsai, C. C.; Connell, G. A. N. *Phys. Rev. Lett.* **1980**, *44*, 273–276. doi:10.1103/physrevlett.44.273
38. Lui, C. H.; Ye, Z.; Ji, C.; Chiu, K.-C.; Chou, C.-T.; Andersen, T. I.; Means-Shively, C.; Anderson, H.; Wu, J.-M.; Kidd, T.; Lee, Y.-H.; He, R. *Phys. Rev. B* **2015**, *91*, 165403. doi:10.1103/physrevb.91.165403
39. Huang, S.; Liang, L.; Ling, X.; Puzos, A. A.; Geoghegan, D. B.; Sumpter, B. G.; Kong, J.; Meunier, V.; Dresselhaus, M. S. *Nano Lett.* **2016**, *16*, 1435–1444. doi:10.1021/acs.nanolett.5b05015
40. Liao, M.; Wei, Z.; Du, L.; Wang, Q.; Tang, J.; Yu, H.; Wu, F.; Zhao, J.; Xu, X.; Han, B.; Liu, K.; Gao, P.; Polcar, T.; Sun, Z.; Shi, D.; Yang, R.; Zhang, G. *Nat. Commun.* **2020**, *11*, 2153. doi:10.1038/s41467-020-16056-4
41. Quan, J.; Linhart, L.; Lin, M.-L.; Lee, D.; Zhu, J.; Wang, C.-Y.; Hsu, W.-T.; Choi, J.; Embley, J.; Young, C.; Taniguchi, T.; Watanabe, K.; Shih, C.-K.; Lai, K.; MacDonald, A. H.; Tan, P.-H.; Libisch, F.; Li, X. *Nat. Mater.* **2021**, *20*, 1100–1105. doi:10.1038/s41563-021-00960-1
42. Tiberj, A.; Rubio-Roy, M.; Paillet, M.; Huntzinger, J.-R.; Landois, P.; Mikolasek, M.; Contreras, S.; Sauvajol, J.-L.; Dujardin, E.; Zahab, A.-A. *Sci. Rep.* **2013**, *3*, 2355. doi:10.1038/srep02355
43. Yan, R.; Simpson, J. R.; Bertolazzi, S.; Brivio, J.; Watson, M.; Wu, X.; Kis, A.; Luo, T.; Hight Walker, A. R.; Xing, H. G. *ACS Nano* **2014**, *8*, 986–993. doi:10.1021/nn405826k
44. Melnikova-Kominkova, Z.; Jurkova, K.; Vales, V.; Drogowska-Horná, K.; Frank, O.; Kalbac, M. *Phys. Chem. Chem. Phys.* **2019**, *21*, 25700–25706. doi:10.1039/c9cp04993b
45. Lee, T.; Choi, J.-H.; Ahn, J.-H.; Yoon, Y.-G.; Rho, H. *Appl. Surf. Sci.* **2022**, *579*, 152208. doi:10.1016/j.apsusc.2021.152208

46. Liu, K.; Zhang, L.; Cao, T.; Jin, C.; Qiu, D.; Zhou, Q.; Zettl, A.; Yang, P.; Louie, S. G.; Wang, F. *Nat. Commun.* **2014**, *5*, 4966. doi:10.1038/ncomms5966
47. Rice, C.; Young, R. J.; Zan, R.; Bangert, U.; Wolverson, D.; Georgiou, T.; Jalil, R.; Novoselov, K. S. *Phys. Rev. B* **2013**, *87*, 081307. doi:10.1103/physrevb.87.081307
48. Li, Z.; Lv, Y.; Ren, L.; Li, J.; Kong, L.; Zeng, Y.; Tao, Q.; Wu, R.; Ma, H.; Zhao, B.; Wang, D.; Dang, W.; Chen, K.; Liao, L.; Duan, X.; Duan, X.; Liu, Y. *Nat. Commun.* **2020**, *11*, 1151. doi:10.1038/s41467-020-15023-3
49. Cortijo-Campos, S.; Prieto, C.; De Andrés, A. *Nanomaterials* **2022**, *12*, 1330. doi:10.3390/nano12081330
50. Liu, Q.; Li, L.; Li, Y.; Gao, Z.; Chen, Z.; Lu, J. *J. Phys. Chem. C* **2012**, *116*, 21556–21562. doi:10.1021/jp307124d
51. Debnath, R.; Maity, I.; Biswas, R.; Raghunathan, V.; Jain, M.; Ghosh, A. *Nanoscale* **2020**, *12*, 17272–17280. doi:10.1039/c9nr09897f
52. Mignuzzi, S.; Pollard, A. J.; Bonini, N.; Brennan, B.; Gilmore, I. S.; Pimenta, M. A.; Richards, D.; Roy, D. *Phys. Rev. B* **2015**, *91*, 195411. doi:10.1103/physrevb.91.195411
53. Zhou, X.; Jin, K.; Cong, X.; Tan, Q.; Li, J.; Liu, D.; Luo, J. *J. Colloid Interface Sci.* **2019**, *538*, 159–164. doi:10.1016/j.jcis.2018.11.032
54. Maino, J. L.; Schouten, R.; Umina, P. *J. Appl. Ecol.* **2021**, *58*, 789–800. doi:10.1111/1365-2664.13812

License and Terms

This is an open access article licensed under the terms of the Beilstein-Institut Open Access License Agreement (<https://www.beilstein-journals.org/bjnano/terms>), which is identical to the Creative Commons Attribution 4.0

International License

(<https://creativecommons.org/licenses/by/4.0>). The reuse of material under this license requires that the author(s), source and license are credited. Third-party material in this article could be subject to other licenses (typically indicated in the credit line), and in this case, users are required to obtain permission from the license holder to reuse the material.

The definitive version of this article is the electronic one which can be found at:

<https://doi.org/10.3762/bjnano.15.26>



On the mechanism of piezoresistance in nanocrystalline graphite

Sandeep Kumar^{*1}, Simone Dehm¹ and Ralph Krupke^{1,2,3}

Full Research Paper

Open Access

Address:

¹Institute of Nanotechnology, Karlsruhe Institute of Technology, Kaiserstr. 12, 76131 Karlsruhe, Germany, ²Institute of Quantum Materials and Technologies, Karlsruhe Institute of Technology, Kaiserstr. 12, 76131 Karlsruhe, Germany and ³Institute of Materials Science, Technische Universität Darmstadt, 64287 Darmstadt, Germany

Email:

Sandeep Kumar^{*} - sandeep.kumar@kit.edu

* Corresponding author

Keywords:

grain boundary; nanocrystalline graphene; strain sensor; Raman; tunneling and destruction

Beilstein J. Nanotechnol. **2024**, *15*, 376–384.

<https://doi.org/10.3762/bjnano.15.34>

Received: 23 November 2023

Accepted: 19 March 2024

Published: 08 April 2024

This article is part of the thematic issue "Applications of Raman spectroscopy in the characterization of nanomaterials".

Guest Editor: P. Araujo



© 2024 Kumar et al.; licensee Beilstein-Institut.
License and terms: see end of document.

Abstract

Strain sensors are sensitive to mechanical deformations and enable the detection of strain also within integrated electronics. For flexible displays, the use of a seamlessly integrated strain sensor would be beneficial, and graphene is already in use as a transparent and flexible conductor. However, graphene intrinsically lacks a strong response, and only by engineering defects, such as grain boundaries, one can induce piezoresistivity. Nanocrystalline graphene (NCG), a derivative form of graphene, exhibits a high density of defects in the form of grain boundaries. It holds an advantage over graphene in easily achieving wafer-scale growth with controlled thickness. In this study, we explore the piezoresistivity in thin films of nanocrystalline graphite. Simultaneous measurements of sheet resistance and externally applied strain on NCG placed on polyethylene terephthalate (PET) substrates provide intriguing insights into the underlying mechanism. Raman measurements, in conjunction with strain applied to NCG grown on flexible glass, indicate that the strain is concentrated at the grain boundaries for smaller strain values. For larger strains, mechanisms such as grain rotation and the formation of nanocracks might contribute to the piezoresistive behavior in nanocrystalline graphene.

Introduction

Flexible strain sensors are an important factor in moving from rigid to flexible electronics. Graphene, because of its interesting inherent properties, has found its way in many applications [1-3]. In particular, it is a promising alternative material as a transparent and conductive coating for future flexible electronics. This is because the relative change in resistance of graphene for similar values of applied strain (4%) is just 50%,

which is two orders of magnitude lower than that of the flat-screen material indium titanium oxide (ITO). Indeed, from a theoretical point of view, change in resistance due to strain or piezoresistivity in graphene is expected to be small because the displacement of the Dirac point occurs in continuous k space, and strain-induced lattice distortions do not change the local band structure up to 20% strain [4]. In contrast, because of the

quantized k space in carbon nanotubes, uniaxial strain can induce band opening or closing. Nevertheless, strain-induced resistance modulation in graphene is by far not zero, and reported values for relative resistance changes vary between 0.1% and 50% at 3% strain [5-8]. Although several works report the enhancement of piezoresistance in graphene, it is still unclear which factors influence this property. A theoretical work by Kumar et al. suggested that grain boundaries can affect piezoresistance in graphene [9]. This result seemed unexpected since Dirac particles should undergo Klein tunneling at barriers without adding up to the total resistance. However, the theoretical modeling shows that the modulation of the transport gap under strain is sensitive to the degree of asymmetry of the grain boundaries. While the symmetric grain boundaries remain metallic in the presence of uniaxial strain, the transport gap of the asymmetric semiconducting grain boundaries can be considerably increased in the presence of strain. Hence, the asymmetric metallic grain boundaries undergo a metal–semiconductor transition in the presence of strain. This effect could open a way to utilize grain boundaries in graphene for fabricating highly sensitive transparent strain sensors.

So far, the growth of specific grain boundaries in graphene has not been reported. Also, most research activities aim at the chemical vapor deposition (CVD) synthesis of monocrystalline graphene free of grain boundaries [10-12]. Methods to detect and visualize grain boundaries and dislocations are currently under development [13,14]. This leads to the situation that the role of grain boundaries for graphene-based sensing of strain, pressure, and motion has not been explored and remains unre-

solved [15-18], although in CVD graphene the domain size is typically of the order of a few micrometers [11]. We speculated that if grain boundaries are responsible for piezoresistivity in graphene, then the gauge factor should be enhanced if one reduces the grain size to a few nanometers.

Nanocrystalline graphene (NCG) is graphitic material with a crystal size of nanometers and, therefore, an excellent candidate for piezoresistance devices. Also, wafer-scale synthesis of NCG has already been achieved by Zhang et al. and modified by Riaz and co-workers [19,20]. Thickness-controlled growth of NCG was exploited to utilize it as an efficient broadband photodetector [21,22]. The preliminary work on the piezoresistivity of NCG looked promising but was limited regarding the applied strain because of rigid SiO_2/Si substrates. In this work, we focus on the piezoresistance measurements in NCG at larger strain values. Initially, a two-point bending setup is described, which was constructed in-house and automated using Python. Then, sheet resistance measurements under externally applied strain are discussed. Raman spectroscopy of the NCG under strain is studied, which gives insights into the distribution of strain in the film. Utilizing electrical and optical properties, a mechanism for piezoresistance in NCG is proposed. The work included here is a part of the PhD thesis completed by the first author S. Kumar [23].

Results and Discussion

The two-point bending fixture, which was constructed to impart external strain and simultaneously perform sheet resistance measurements and Raman spectroscopy, is shown in Figure 1a.

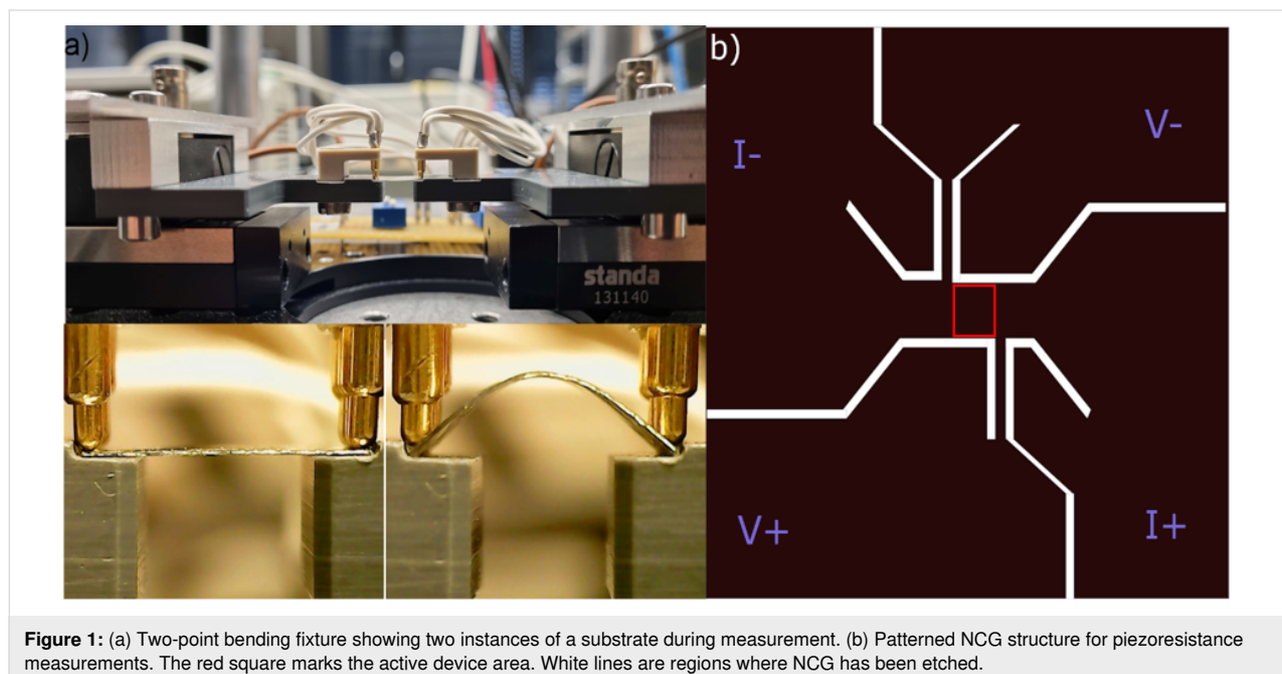
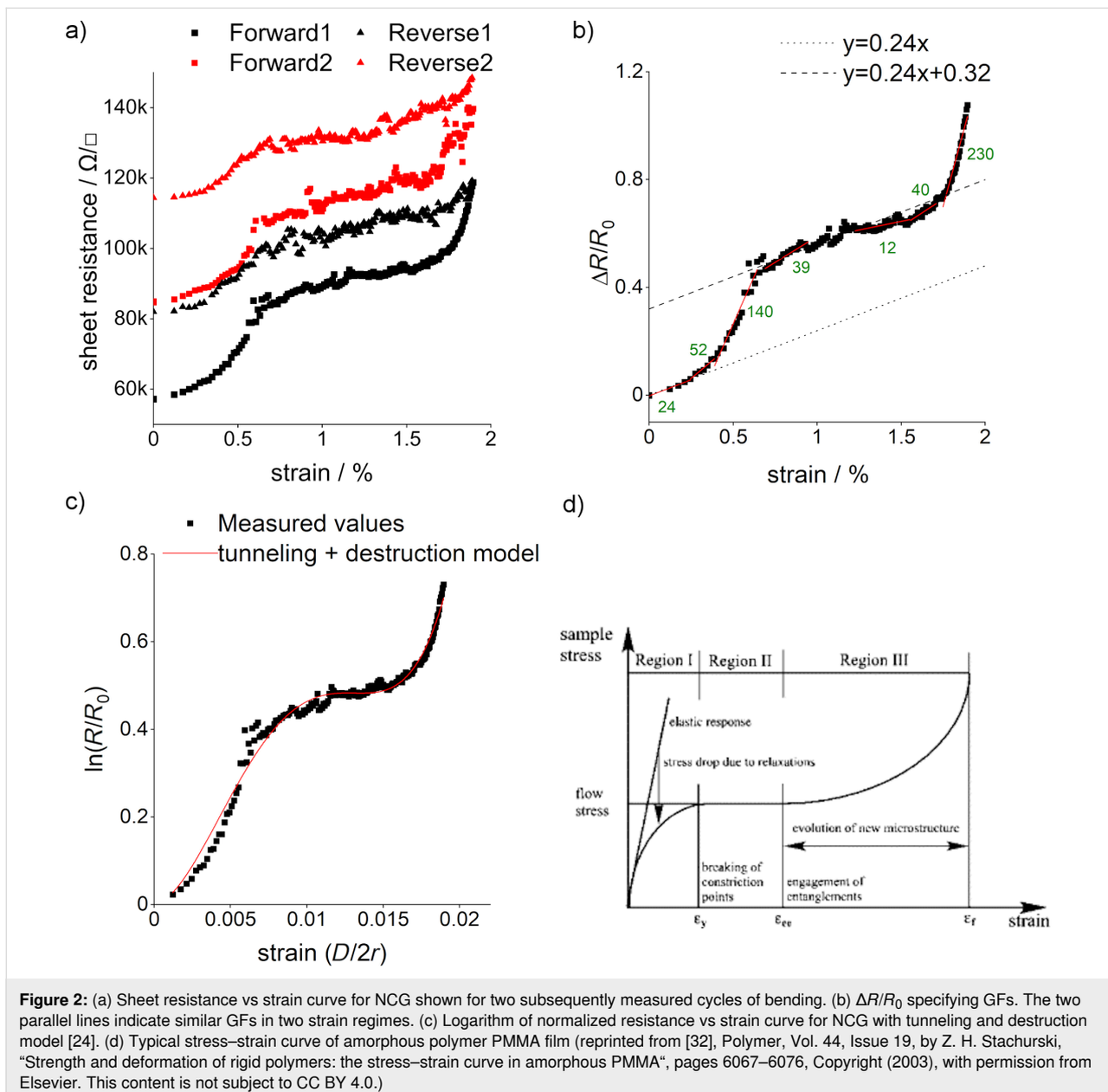


Figure 1: (a) Two-point bending fixture showing two instances of a substrate during measurement. (b) Patterned NCG structure for piezoresistance measurements. The red square marks the active device area. White lines are regions where NCG has been etched.

Two instances of zero and non-zero strain are also depicted. The setup has two stepper motors acquired from Standa Inc. The contacts on the NCG were made by gold-coated spring pressure contacts, which were then connected to a BNC connector and a Keithley 2636A device. The substrate holder and contacts holder were machined and attached to the stepper motor as shown in Figure 1a. A detailed description of the setup has been given by Kumar [23]. The complete setup was automated via self-programmed Python code. To completely eliminate any strain-induced changes in the contacts, the NCG was patterned such that the NCG itself is used as a contacting electrode (shown in Figure 1b). The area marked with a red square in Figure 1b (2 mm × 2 mm) is the active device area for sheet

resistance measurements on a substrate of 10 mm × 10 mm area. Thin NCG constrictions at the end of the active device area were used to measure the potential drop across the device area. The measurements were done in constant current mode, and the voltage drop across the squared central area was measured at each strain value.

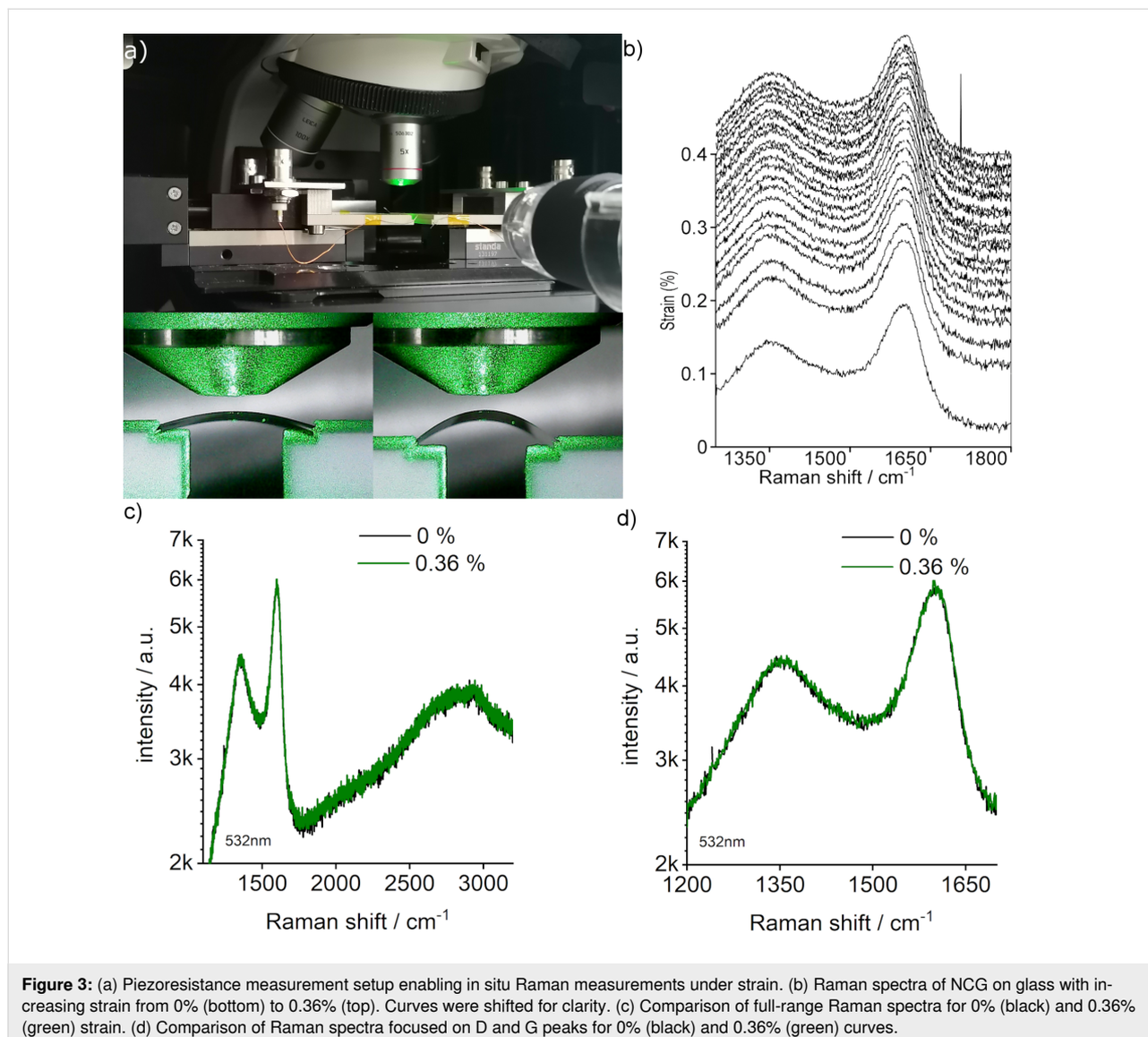
NCG was grown by spin coating S1805 at 4000 rpm; subsequently, it was transferred onto a 100 μm thick PET substrate. For more details please see the Experimental section. Measured piezoresistance curves (with forward and reverse sweep) for a 5 nm thick patterned NCG film on the PET substrate are shown in Figure 2a. Sheet resistance values are plotted against tensile



strain up to 2%, which is an order of magnitude larger than in our previous work (max 0.1%) [19]. We could reproduce the previously observed gauge factor (GF) of ca. 24 at very low strain (<0.3%); however, in the extended-strain region we observe now a super-linear behavior, which evolves into a linear region from 0.7% to 1.6% strain, before entering a super-linear regime beyond 1.6% strain. The GF values from the plot of normalized change in resistance vs strain (Figure 2b) also depict how the rate of resistance change with strain drops and then increases again. Interestingly, the GF is similar below 0.3% and in the region between 0.7% and 1.6% strain, indicating a similar origin of piezoresistance (shown by two parallel lines in Figure 2b). The overall shape of the curves is reproducible and shown here for two strain cycles. However, a hysteresis is observed between forward and reverse sweeps, indicating that

structural changes in the films occur, which are in part irreversible.

To gain insights into the strain distribution in the strained NCG, we performed in situ Raman measurements with strain as shown in Figure 3a. The flexible glass was preferred for the Raman measurements because the spectra of NCG cannot be resolved on PET as a result of a strong Raman signal of the substrate itself. The 50 μm thick flexible glass was acquired from Schott. The glass loses its flexibility at 600 $^{\circ}\text{C}$ and also in water [23]. To keep the flexibility, the NCG film was grown on both sides of the glass substrate. There are three reasons for that. First, the negative thermal expansion coefficient of NCG prevents the release of stress initially present in the glass [25]. Second, the film protects the glass from any corrosion from water if the transfer is required on glass in an aqueous medium



[26]. Third, NCG fills the cracks present at the edges during spin coating the polymer and inhibits their propagation during the bending of the substrate.

Raman spectroscopy is a powerful method to detect strain in graphene, which can be determined from the analysis of the peak position of the 2D and G modes [27]. For the Raman measurements, the bending setup was installed in a Renishaw inVia Raman microscope, operated at 532 nm excitation wavelength (Figure 3a) with a 100× objective. Concado et al. [28] reported the general equation for calculating the crystallite size based on the full width at half maximum (FWHM) and intensity ratios of D and G peaks. Similar to our previous work [19], we use the FWHM of D and G peaks here at zero strain to calculate the crystallite size. The G peak gives a crystallite size of 2–3 nm, and the D peak corresponds to a crystallite size of 4–5 nm. TEM studies also done in our previous work on NCG gave an average domain size of 3 nm. We therefore report a grain size range of 2–5 nm for the NCG film synthesized for this study.

Figure 3b shows the Raman spectra as a waterfall plot for strain values up to 0.36%. Figure 3c,d shows a comparison between the spectra taken at 0% and 0.36% strain. Further measurements with increased strain could not be taken due to the failure of the glass substrate. Interestingly, no prominent changes in peak positions, widths, or intensity could be detected, inferring that the strain within the grains remains constant even though the externally applied strain increased to ca. 0.4%.

In an attempt to model piezoresistance in NCG, we have used the tunneling + destruction model for composite materials [24]:

$$\ln \frac{R}{R_0} = \ln(1 + \epsilon) + \alpha\epsilon + \beta\epsilon^2 + \gamma\epsilon^3 + \delta\epsilon^4.$$

The model with five free parameters was fitted to the data as shown in Figure 2c, and the fit parameters are given in Table 1. The model was initially given for a matrix in which conducting particles are dispersed in a polymer matrix and are separated by tunnel junctions. In this model, the conductivity in the film is determined by the number of conductive paths, N , and the tunneling distance, d . The model has been used to explain the piezoresistance for several composite materials [29,30].

Zhao et al. [24] used the model to explain the piezoresistance in nanographene films, although the material is comparable to ours and not a composite material in the original sense. NCG can be considered as a matrix of grains and grain boundaries (GBs), where the grains are separated by the GBs and have different resistivities [31]. Hence, the tunneling + destruction model might indeed be an appropriate physical representation of the

Table 1: Value of extracted parameters by fitting from Figure 2c.

Parameter	Value
α	5
β	$1.55 \times 10^4 \pm 1.8 \times 10^2$
γ	$-1.63 \times 10^6 \pm 2.5 \times 10^4$
δ	$4.6 \times 10^7 \pm 8.6 \times 10^5$

nano/microstructure of NCG. The model explains that at lower strain values, only the tunneling distance d increases, but N remains constant. Whereas at larger strain values, d and N both change; therefore, the GF increases at those values. Our Raman measurements indicate that at least up to 0.4% strain, no strain is experienced by the grains, which would mean that all the strain energy provided externally ends up at GBs leading to movements of dislocations at GBs or fractures at GBs. Fitting the tunneling + destruction model to our data, we find that the initial tunneling distance has a value of 3.6 nm, which is comparable to the nanographene films fabricated by Zhao et al. (3.4 nm) [24]. The destruction of conduction channels as part of the tunneling + destruction model then takes place at higher strain values and eventually leads to partial irreversibility, which can be observed as an offset between the first and second trace in Figure 2a. Nevertheless, the overall shape of the second curve is similar to the first one. Figure 2d shows a typical stress vs strain curve for a polymer film [32]. The trend of the curve looks similar to the resistance vs strain curve for NCG in Figure 2a. Since polymers are insulators, literature on resistance vs strain for such films does not exist. However, it depicts how the stress drops because of strain relaxation in such films. Since resistance is directly proportional to strain in the tunneling + destruction model, one can think that the resistance of such films would also drop. Based on strain relaxations, therefore, one can correlate NCG and a polymer material and potentially give insight into how NCG behaves in the plateau region of the resistance vs strain curve.

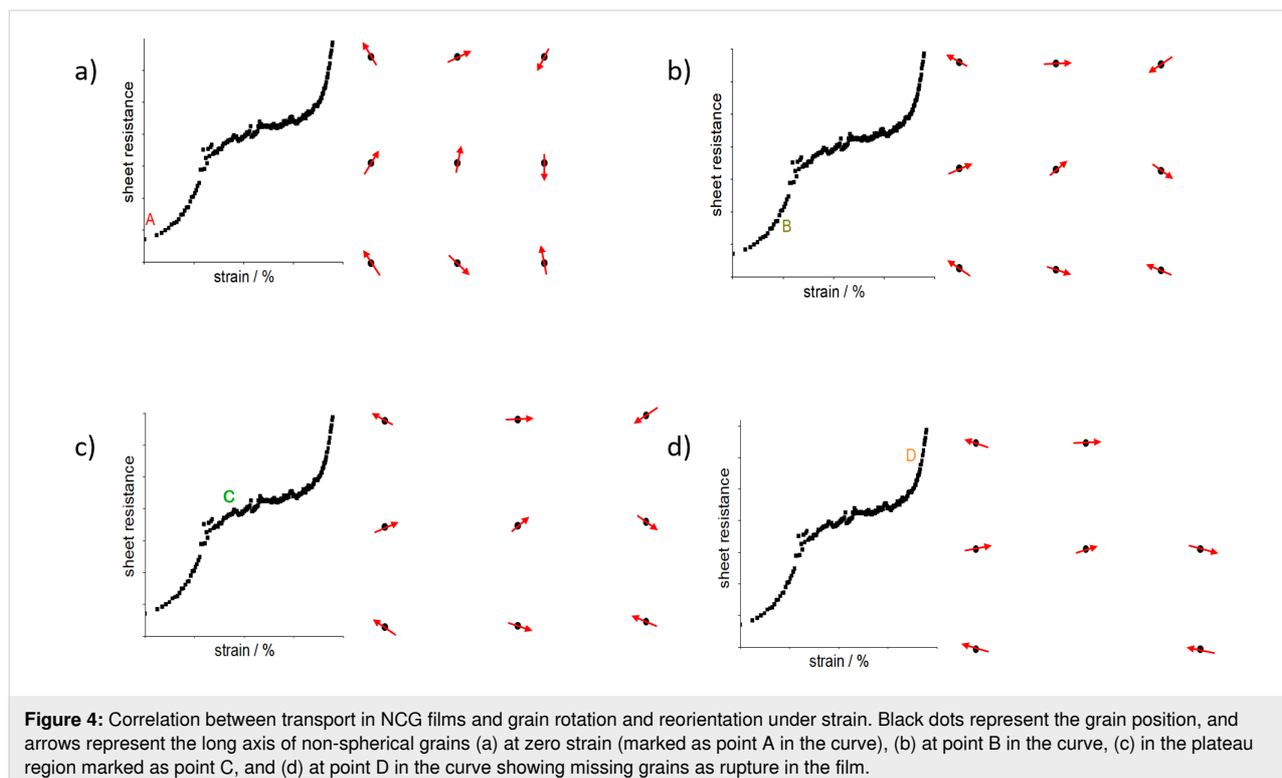
It is important to note that many works that have reported piezoresistance in NCG have recorded data at comparably low strain values and have not observed the plateau-like region as reported here, where the gauge factor is similar to the gauge factor at very low strain [24,33]. A plateau-like region has neither been observed in nanocrystalline graphite [33], amorphous carbon films [34], nor in metallic films [35]. The mechanism that leads to an increase of resistance in amorphous carbon and gold films at large strain is crack formation. Also, in NCG, which is full of GBs and defects, crack formation and propagation have to be considered [36]. Assuming nanocrack formation at the GBs, we could understand the entire piezoresistance curve in the following way. The increase in resistance at

lower strain (0.3%) would be determined by the piezoresistance of the GBs, as studied by Kumar and co-workers [9]. Beyond this strain value, the resistance of the film increases rapidly, which can be understood by nanocrack formation at certain GBs [37]. This phenomenon occurs up to 0.7% strain and then stops. The reason could be that the remaining GBs are stronger than the ones that fractured between 0.3% and 0.7% strain. The plateau after 0.7% strain indicates that although the strain is increased, no crack formation occurs. In addition, the slopes of the resistance vs strain curve at smaller strain and the plateau region are similar (Figure 2b), indicating a similar piezoresistance mechanism in both strain regions. This can be understood by strain relaxation occurring at GBs by crack formation and bond movement, as well as reformation and changes in the microstructure of the film [37]. After the plateau (strain > 1.6%), the resistance once again increases, and a new set of nanocracks start to form at different GBs. The second cycle of the resistance vs strain measurement (Figure 2a, red curve) starts from a resistance value that is equivalent to the resistance at 0.7% strain in the first cycle. This indicates that the nanocracks formed in the NCG film between 0.3% to 0.7% strain are non-reversible cracks. However, the cracks formed between 1.6% and 2.0% strain are partially reversible, and the resistance values are almost recovered. This is possible through bond reformations by formation of pentagon–heptagon pairs due to their low formation energy [38,39]. In addition, the formation of nanocracks appears not so pronounced in the second cycle, as

can be seen from the change in resistance between 0.3% and 0.7% and between 1.6% and 2.0% strain, and is likely due to cracks already formed in the first cycle.

Another process that might be considered for strain relaxation is grain rotation as shown in Figure 4. The rotation of grains during the straining of metal films has been studied extensively [40,41]. The effect is prominent for metal films with smaller grain sizes and diminishes with larger grains. The mechanism has been proposed as a cause of plastic deformation in metals. Wang et al. showed for a platinum nanocrystalline film [42] that the rotation due to strain at room temperature does not occur because of cross-grain gliding, GB sliding, or diffusional creep processes. The rotation occurs because of a change in the content of GB dislocations, which can change the GB angle between the grains (Frank–Bilby equation).

We cannot completely exclude that GB rotation occurs to some extent also in strained nanocrystalline graphene. Figure 4 shows a schematic of a NCG film under strain correlated with the transport of the film. The black dots represent the grain position, and the arrows represent the long axis of non-spherical grains. At the beginning, marked as point A on the curve, the grains are randomly oriented. As the strain is applied, the grains start to move apart, which is visible as an increase in resistance values between 0% and 0.3% strain [24], also confirmed by Raman measurements under strain (Figure 3b–d). At 0.3% to



0.7% strain, grain rotation and irreversible changes in the microstructure occur. This is seen by a sharper increase in resistance and a larger GF value, which corresponds to point B in Figure 4b showing grains moving apart in combination with grain rotation. At the plateau region (0.7%–1.6%), as explained before, the slope of the resistance vs strain curve is equivalent to the lower strain region (0%–0.3%) indicating a similar piezoresistance mechanism of grains moving apart and increasing tunneling distance. This is shown as point C in the transport in Figure 4c, where the grains are locked and cannot rotate, and the increase in resistance only occurs because of increased distance between grains. Above 1.6% strain, a sharper increase in resistance indicates again grain rotation and reorientation, and fracture in the film shown by missing black dots and arrows in Figure 4d, corresponding to point D in the transport curve. The processes of grain movements (increase in distance between grains and rotation) would repeat if the strain values are increased further until the fracture of the film. The process of bond rotation and reformation is known in NCG films for the relaxation of stress at GBs [37]. When the application of strain is reversed, irreversible changes occurring in the film by grain rotation results in a permanent increase in initial resistance seen by a hysteresis (black and red curves in Figure 2a) and an offset in the second cycle (red curve in Figure 2a) [43]. Although the processes are different, there is a competition between bond breaking and rotation at certain strain values; the kinetically favorable process occurs in alternating sequences related to thermal and stress fluctuations inducing nanocrack formation [37,44]. Yang et al. [45] have shown a simulation of the stress vs strain behavior in NCG films at different temperatures and strain rates. Interestingly, the curve looks similar to the resistance vs strain curve in this work. A deviation from the linear behavior into a plateau is observed at larger strain, owing to plastic deformations of the NCG film. However, a sharp increase in stress after the plateau region is not observed. This can be understood by stress relaxation due to fractures in the film at higher strain, which is visible in resistance vs strain curves as a sharp increase in the resistance. As a last comment, Zhao et al. [24] reported that as they reduced the grain size from 25 to 8 nm, the GF increased from 11 to 600. Also, Simionescu et al. [33] reported a varying GF (50–250) for a strain range of 0%–1%. In this work, NCG with lower grain sizes has been obtained; however, the GF does not appear to further increase and remains comparable to the values of previously reported works. A comparison is tabulated in Table 2.

Conclusion

This study endeavors to further the understanding of the piezoresistance mechanism in NCG, employing a two-point bending setup to apply controlled strain. The strained NCG was analyzed electrically and optically, revealing three regimes in

Table 2: Comparison of grain sizes of NCG with the corresponding GFs.

Grain size (nm)	Gauge factor (GF)	Strain range (%)	Reference
25	11	0 to 1	[24]
8	600	0 to 1	[24]
9	50–250	0 to 1	[33]
2–5	23	0 to 0.1	[19]
2–5	24–140	0 to 2	this work

the sheet resistance vs strain curve. Examination of the results from optical and electrical measurements suggests that in the lower strain regime, the grains experience negligible effects, while the majority of strain is concentrated at the grain boundaries. Consequently, non-reversible cracks form at GBs. The second regime exhibits a superlinear dependence of sheet resistance on strain, indicating potential grain rotation and bond reformation, leading to a modified nano/microstructure. In the larger-strain regime, an exponential increase in sheet resistance vs strain signifies further partially reversible crack formation. To enhance understanding, a tunneling + destruction model was fitted, and parameters were extracted. While the paper offers an overview of piezoresistance in NCG, a more in-depth study is imperative for a complete comprehension of the system's complexity. In situ FTIR measurements could provide additional insights into changes in doping and defects with strain.

Experimental

Piezoresistance measurements

NCG was synthesized on a 300 nm SiO₂/Si substrate by spin coating S1805 (1:10 dilution with propylene glycol methyl ether acetate, PGMEA) at 4000 rpm. The spin-coated Si/SiO₂ substrate was loaded in a vacuum furnace and annealed at 600 °C for 10 h at 10^{−6} mbar. The measured thickness of the grown film was ca. 5 nm. The NCG film was then transferred onto a 100 μm thick PET substrate. For the transfer process, first, the NCG film on SiO₂/Si was coated with 200 nm thick PMMA and put into 5 M NaOH solution at 80 °C. The NCG/PMMA film floats on the surface after the etching of SiO₂. Using a clean glass wafer, the NCG/PMMA film was transferred from the NaOH solution to a clean water beaker and allowed to float on the top. The cleaning was repeated three times to ensure the no residues of NaOH remained on the NCG film. The film was then removed from the water using a PET substrate. After that, the substrate was left in air for drying. Next, a drop of PMMA was dripped on top of the film and allowed to spread and dry. This has been shown to be helpful in removing wrinkles formed during the transfer process [46]. The NCG film on the PET substrate was then patterned in the structure shown in Figure 1b

using e-beam lithography. There were no metal films deposited on NCG, and the electrical contact was made between gold spring contacts and NCG directly. For Raman measurements, S1805 (1:10 dilution with PGMEA) was spin-coated on both sides of the flexible glass substrate at 4000 rpm to grow NCG on both sides of the glass. The substrate was then loaded into the vacuum furnace and treated similarly.

Raman measurements

Raman measurements were done using a 100× objective at 0.6 mW laser power for 60 s integration time for each measurement. The same area on the NCG film was focused as to monitor and compare any changes occurring during straining the film.

ORCID® iDs

Sandeep Kumar - <https://orcid.org/0000-0003-0991-9438>

Simone Dehm - <https://orcid.org/0000-0002-3431-4732>

Ralph Krupke - <https://orcid.org/0000-0001-8427-8592>

References

- Choi, J. H.; Lee, J.; Byeon, M.; Hong, T. E.; Park, H.; Lee, C. Y. *ACS Appl. Nano Mater.* **2020**, *3*, 2257–2265. doi:10.1021/acsnm.9b02378
- Fu, W.; Nef, C.; Knopfmacher, O.; Tarasov, A.; Weiss, M.; Calame, M.; Schönenberger, C. *Nano Lett.* **2011**, *11*, 3597–3600. doi:10.1021/nl201332c
- Yavari, F.; Castillo, E.; Gullapalli, H.; Ajayan, P. M.; Koratkar, N. *Appl. Phys. Lett.* **2012**, *100*, 203120. doi:10.1063/1.4720074
- Pereira, V. M.; Castro Neto, A. H.; Peres, N. M. R. *Phys. Rev. B* **2009**, *80*, 045401. doi:10.1103/physrevb.80.045401
- Bae, S.; Kim, H.; Lee, Y.; Xu, X.; Park, J.-S.; Zheng, Y.; Balakrishnan, J.; Lei, T.; Ri Kim, H.; Song, Y. I.; Kim, Y.-J.; Kim, K. S.; Özyilmaz, B.; Ahn, J.-H.; Hong, B. H.; Iijima, S. *Nat. Nanotechnol.* **2010**, *5*, 574–578. doi:10.1038/nnano.2010.132
- Bae, S.-H.; Lee, Y.; Sharma, B. K.; Lee, H.-J.; Kim, J.-H.; Ahn, J.-H. *Carbon* **2013**, *51*, 236–242. doi:10.1016/j.carbon.2012.08.048
- Fu, X.-W.; Liao, Z.-M.; Zhou, J.-X.; Zhou, Y.-B.; Wu, H.-C.; Zhang, R.; Jing, G.; Xu, J.; Wu, X.; Guo, W.; Yu, D. *Appl. Phys. Lett.* **2011**, *99*, 213107. doi:10.1063/1.3663969
- Li, X.; Zhang, R.; Yu, W.; Wang, K.; Wei, J.; Wu, D.; Cao, A.; Li, Z.; Cheng, Y.; Zheng, Q.; Ruoff, R. S.; Zhu, H. *Sci. Rep.* **2012**, *2*, 870. doi:10.1038/srep00870
- Kumar, S. B.; Guo, J. *Nano Lett.* **2012**, *12*, 1362–1366. doi:10.1021/nl203968j
- Lee, J.-H.; Lee, E. K.; Joo, W.-J.; Jang, Y.; Kim, B.-S.; Lim, J. Y.; Choi, S.-H.; Ahn, S. J.; Ahn, J. R.; Park, M.-H.; Yang, C.-W.; Choi, B. L.; Hwang, S.-W.; Whang, D. *Science* **2014**, *344*, 286–289. doi:10.1126/science.1252268
- Bonaccorso, F.; Lombardo, A.; Hasan, T.; Sun, Z.; Colombo, L.; Ferrari, A. C. *Mater. Today* **2012**, *15*, 564–589. doi:10.1016/s1369-7021(13)70014-2
- Li, X.; Magnuson, C. W.; Venugopal, A.; Tromp, R. M.; Hannon, J. B.; Vogel, E. M.; Colombo, L.; Ruoff, R. S. *J. Am. Chem. Soc.* **2011**, *133*, 2816–2819. doi:10.1021/ja109793s
- Butz, B.; Dolle, C.; Niekkel, F.; Weber, K.; Waldmann, D.; Weber, H. B.; Meyer, B.; Spiecker, E. *Nature* **2014**, *505*, 533–537. doi:10.1038/nature12780
- Duong, D. L.; Han, G. H.; Lee, S. M.; Gunes, F.; Kim, E. S.; Kim, S. T.; Kim, H.; Ta, Q. H.; So, K. P.; Yoon, S. J.; Chae, S. J.; Jo, Y. W.; Park, M. H.; Chae, S. H.; Lim, S. C.; Choi, J. Y.; Lee, Y. H. *Nature* **2012**, *490*, 235–239. doi:10.1038/nature11562
- Smith, A. D.; Niklaus, F.; Pausa, A.; Vaziri, S.; Fischer, A. C.; Sterner, M.; Forsberg, F.; Delin, A.; Esseni, D.; Palestri, P.; Östling, M.; Lemme, M. C. *Nano Lett.* **2013**, *13*, 3237–3242. doi:10.1021/nl401352k
- Zhu, S.-E.; Krishna Ghatkesar, M.; Zhang, C.; Janssen, G. C. A. M. *Appl. Phys. Lett.* **2013**, *102*, 161904. doi:10.1063/1.4802799
- Hosseinzadegan, H.; Todd, C.; Lal, A.; Pandey, M.; Levendorf, M.; Park, J. Graphene has ultra high piezoresistive gauge factor. In *2012 IEEE 25th International Conference on Micro Electro Mechanical Systems (MEMS)*, Paris, France, Jan 29–Feb 2, 2012; IEEE; pp 611–614. doi:10.1109/memsys.2012.6170262
- Smith, A. D.; Vaziri, S.; Niklaus, F.; Fischer, A. C.; Sterner, M.; Delin, A.; Östling, M.; Lemme, M. C. *Solid-State Electron.* **2013**, *88*, 89–94. doi:10.1016/j.sse.2013.04.019
- Riaz, A.; Pyatkov, F.; Alam, A.; Dehm, S.; Felten, A.; Chakravadhanula, V. S. K.; Flavel, B. S.; Kübel, C.; Lemmer, U.; Krupke, R. *Nanotechnology* **2015**, *26*, 325202. doi:10.1088/0957-4484/26/32/325202
- Zhang, Z.; Ge, B.; Guo, Y.; Tang, D.; Wang, X.; Wang, F. *Chem. Commun.* **2013**, *49*, 2789–2791. doi:10.1039/c3cc00089c
- Peyyety, N. A.; Kumar, S.; Li, M.-K.; Dehm, S.; Krupke, R. *ACS Appl. Mater. Interfaces* **2022**, *14*, 9525–9534. doi:10.1021/acsmi.1c24306
- Parmar, D.; Dehm, S.; Peyyety, N. A.; Kumar, S.; Krupke, R. *Adv. Sens. Res.* **2024**, *3*, 2300134. doi:10.1002/adrs.202300134
- Kumar, S. *Nanocarbon Devices and Sensors*. Ph.D. Thesis, TU Darmstadt, Germany, 2022. doi:10.26083/tuprints-00021384
- Zhao, J.; Wang, G.; Yang, R.; Lu, X.; Cheng, M.; He, C.; Xie, G.; Meng, J.; Shi, D.; Zhang, G. *ACS Nano* **2015**, *9*, 1622–1629. doi:10.1021/nn506341u
- Yoon, D.; Son, Y.-W.; Cheong, H. *Nano Lett.* **2011**, *11*, 3227–3231. doi:10.1021/nl201488g
- Singh Raman, R. K.; Tiwari, A. *JOM* **2014**, *66*, 637–642. doi:10.1007/s11837-014-0921-3
- Lee, J. E.; Ahn, G.; Shim, J.; Lee, Y. S.; Ryu, S. *Nat. Commun.* **2012**, *3*, 1024. doi:10.1038/ncomms2022
- Cançado, L. G.; Takai, K.; Enoki, T.; Endo, M.; Kim, Y. A.; Mizusaki, H.; Jorio, A.; Coelho, L. N.; Magalhães-Paniago, R.; Pimenta, M. A. *Appl. Phys. Lett.* **2006**, *88*, 163106. doi:10.1063/1.2196057
- Al-solamy, F. R.; Al-Ghamdi, A. A.; Mahmoud, W. E. *Polym. Adv. Technol.* **2012**, *23*, 478–482. doi:10.1002/pat.1902
- Zhang, X.-W.; Pan, Y.; Zheng, Q.; Yi, X.-S. *J. Polym. Sci., Part B: Polym. Phys.* **2000**, *38*, 2739–2749. doi:10.1002/1099-0488(20001101)38:21<2739::aid-polb40>3.0.co;2-o
- Zhao, T.; Xu, C.; Ma, W.; Liu, Z.; Zhou, T.; Liu, Z.; Feng, S.; Zhu, M.; Kang, N.; Sun, D.-M.; Cheng, H.-M.; Ren, W. *Nat. Commun.* **2019**, *10*, 4854. doi:10.1038/s41467-019-12662-z
- Stachurski, Z. H. *Polymer* **2003**, *44*, 6067–6076. doi:10.1016/s0032-3861(03)00554-8
- Simionescu, O.-G.; Pachi, C.; Ionescu, O.; Dumbrăvescu, N.; Buiu, O.; Popa, R. C.; Avram, A.; Dinescu, G. *Rev. Adv. Mater. Sci.* **2020**, *59*, 306–313. doi:10.1515/rams-2020-0031
- Leterrier, Y.; Pinyol, A.; Rougier, L.; Waller, J. H.; Månson, J.-A. E. *J. Appl. Phys.* **2009**, *106*, 113508. doi:10.1063/1.3266001

35. Cho, C.; Kang, P.; Taqieddin, A.; Jing, Y.; Yong, K.; Kim, J. M.; Haque, M. F.; Aluru, N. R.; Nam, S. *Nat. Electron.* **2021**, *4*, 126–133. doi:10.1038/s41928-021-00538-4
36. Shekhawat, A.; Ritchie, R. O. *Nat. Commun.* **2016**, *7*, 10546. doi:10.1038/ncomms10546
37. Yang, Z.; Huang, Y.; Ma, F.; Sun, Y.; Xu, K.; Chu, P. K. *Mater. Sci. Eng., B* **2015**, *198*, 95–101. doi:10.1016/j.mseb.2015.03.019
38. Zazyev, O. V. *Solid State Commun.* **2012**, *152*, 1431–1436. doi:10.1016/j.ssc.2012.04.045
39. Malola, S.; Häkkinen, H.; Koskinen, P. *Phys. Rev. B* **2010**, *81*, 165447. doi:10.1103/physrevb.81.165447
40. Argon, A. S.; Yip, S. *Philos. Mag. Lett.* **2006**, *86*, 713–720. doi:10.1080/09500830600986091
41. Yamakov, V.; Wolf, D.; Phillpot, S. R.; Mukherjee, A. K.; Gleiter, H. *Nat. Mater.* **2004**, *3*, 43–47. doi:10.1038/nmat1035
42. Wang, L.; Teng, J.; Liu, P.; Hirata, A.; Ma, E.; Zhang, Z.; Chen, M.; Han, X. *Nat. Commun.* **2014**, *5*, 4402. doi:10.1038/ncomms5402
43. Yang, Z.; Huang, Y.; Ma, F.; Miao, Y.; Bao, H.; Xu, K.; Chu, P. K. *RSC Adv.* **2016**, *6*, 60856–60861. doi:10.1039/c6ra05167g
44. Ovid'ko, I. A.; Sheinerman, A. G. *J. Phys. D: Appl. Phys.* **2013**, *46*, 345305. doi:10.1088/0022-3727/46/34/345305
45. Yang, Z.; Huang, Y.; Ma, F.; Sun, Y.; Xu, K.; Chu, P. K. *Eur. Phys. J. B* **2015**, *88*, 135. doi:10.1140/epjb/e2015-50850-x
46. Li, X.; Zhu, Y.; Cai, W.; Borysiak, M.; Han, B.; Chen, D.; Piner, R. D.; Colombo, L.; Ruoff, R. S. *Nano Lett.* **2009**, *9*, 4359–4363. doi:10.1021/nl902623y

License and Terms

This is an open access article licensed under the terms of the Beilstein-Institut Open Access License Agreement (<https://www.beilstein-journals.org/bjnano/terms>), which is identical to the Creative Commons Attribution 4.0 International License (<https://creativecommons.org/licenses/by/4.0>). The reuse of material under this license requires that the author(s), source and license are credited. Third-party material in this article could be subject to other licenses (typically indicated in the credit line), and in this case, users are required to obtain permission from the license holder to reuse the material.

The definitive version of this article is the electronic one which can be found at:
<https://doi.org/10.3762/bjnano.15.34>



Further insights into the thermodynamics of linear carbon chains for temperatures ranging from 13 to 300 K

Alexandre Rocha Paschoal^{*1,2}, Thiago Alves de Moura^{1,3},
Juan S. Rodríguez-Hernández¹, Carlos William de Araujo Paschoal¹, Yoong Ahm Kim⁴,
Morinobu Endo⁵ and Paulo T. Araujo^{*2}

Full Research Paper

[Open Access](#)

Address:

¹Department of Physics, Federal University of Ceara, 60440-900 Fortaleza, Ceara, Brazil, ²Department of Physics and Astronomy, University of Alabama, Tuscaloosa, Alabama 35487, USA, ³Departamento de Ensino, Instituto Federal de Educação, Ciência e Tecnologia do Ceará (IFCE), 62580-000, Acarau, Ceara, Brazil, ⁴Department of Polymer Engineering, School of Polymer Science and Engineering, and Alan G. MacDiarmid Energy Research Institute, Chonnam National University, 77 Yongbong-ro, Buk-gu, Gwangju 61186, Republic of Korea and ⁵Faculty of Engineering, Shinshu University, 4-17-1 Wakasato, Nagano-shi 380-8553, Japan

Email:

Alexandre Rocha Paschoal^{*} - paschoal@fisica.ufc.br;
Paulo T. Araujo^{*} - paulo.t.araujo@ua.edu

* Corresponding author

Keywords:

carbon nanotubes; Debye model; Grüneisen parameter; linear carbon chains; Raman spectroscopy

Beilstein J. Nanotechnol. **2025**, *16*, 1818–1825.

<https://doi.org/10.3762/bjnano.16.125>

Received: 14 June 2025

Accepted: 09 September 2025

Published: 20 October 2025

This article is part of the thematic issue "Applications of Raman spectroscopy in the characterization of nanomaterials".

Associate Editor: P. Ayala



© 2025 Paschoal et al.; licensee Beilstein-Institut.
License and terms: see end of document.

Abstract

It was recently shown that small bundles of linear carbon chains (LCC) encapsulated by double- and multi-wall carbon nanotubes (LCC@DWCNT and LCC@MWCNT, respectively) behave as Debye's materials for temperatures as high as 293 K with an estimate that such materials could still withstand such characteristics for even higher temperatures (≈ 700 K). Using the Debye model, thermodynamic observables (internal energy, coefficient of linear thermal expansion, specific heat, thermal strain, and Grüneisen parameter at constant pressure) were empirically determined for the first time in the range of temperatures $70 < T < 293$ K. These observables were all correlated with the C-band frequency (ω_{LCC}) dependence on the temperature (T) and its first and second derivatives with relation to T , $d\omega_{\text{LCC}}/dT$, and $d^2\omega_{\text{LCC}}/dT^2$. The C-band is a Raman spectroscopic signature for LCC, which is not only temperature-dependent but also dependent on the number of carbon atoms (N) constituting the LCC. In this present study, we extend these findings to temperatures ranging from $13 < T < 293$ K, which provide more accurate values for both $d\omega_{\text{LCC}}/dT$ and $d^2\omega_{\text{LCC}}/dT^2$. The corrected values of these derivatives affect the Grüneisen parameters associated with the LCC, even though the other associated thermodynamic parameters remain essentially unchanged. Our measurements were performed in both isolated and small bundles of LCC@MWCNT, which allowed us to demonstrate that small bundles or isolated environments do not seem to influence the vibrational and thermodynamic properties measured.

Introduction

Phonons, their mutual interactions (ph–ph interactions), and their interactions with electrons (e–ph interactions) play fundamental roles in how materials respond to electric (e.g., difference of potentials), thermal (e.g., temperature gradients), and mechanical (e.g., pressure variations) stimuli [1-23]. These responses are directly connected with electronic and transport properties, which in turn depend on the equilibrium between emission and absorption of phonons, and gain and loss of energy of carriers [1,2,10,17,24-28]. The phonon lifetime as well as the selection rules behind ph–ph and e–ph interactions determine the efficiency of such phonon emission and absorption [1,2,10,17,24-28]. Phonons need to be in an excited state to be emitted or absorbed. Once they decay to their ground state, they become unavailable. This decay process is often accomplished via three-phonon processes (called the Klemens' channel) and via four-phonon processes [1-3,6,7,12,13,16,24]. It is widely known that pressure (P)- and temperature (T)-dependent phenomena are ruled by anharmonic ph–ph interactions, which are also driven by three- and four-phonon processes, and by e–ph interactions [1-29].

Therefore, phonon assignments in materials as well as the understanding of how such phonons relate to thermal and mechanical properties of the materials become of fundamental importance [1-30]. One important point to keep in mind is that ph–ph and e–ph interactions are also very susceptible to the dimensionality of the materials, and for one-dimensional (1D) materials, the selection rules behind such interactions are rather restricted [13,31-33]. These interactions are all quantum-related phenomena, and their ineffectiveness allows thermal and mechanical properties of materials to be described by semi-classical theories (such as the Debye's theory that describes the behavior of materials with T), and their phonon frequencies might be directly connected with relevant parameters such as the Young's modulus, the Grüneisen parameter, and thermal expansion coefficient [29,30]. This is the case with linear atomic chains constituted of carbon atoms [29,30].

Linear carbon chains (LCC) are 1D systems that are classified into two categories: polyynes (displaying alternating triple and single bonds between constituent carbon atoms) and cumulenes (displaying only double bonds between constituent carbon atoms) [31,34-40]. Cumulenes are metallic systems that, due to Peierls transition, are more unstable than polyynes, which present insulating properties with bandgaps whose sizes are dependent on the number of carbon atoms (N) constituting the chains [31,34-40]. The unique properties associated with LCC have attracted a great deal of attention in the scientific community. They are structures that present unique anharmonic behaviors [29,30,41,42], and they are claimed to possess one of the

largest mechanical resistances among materials (including other carbon allotropic versions like graphene or nanotubes) [29,43-49], in addition to presenting unique conductive properties that place them ahead as ideal candidates for future developments in nanoelectronics [43-49]. Moreover, due to its simplicity, LCC are like textbook problems in which many simple, but powerful, theories can be tested [29-31].

Until recently, many challenging questions regarding LCC stability have been raised [39,50-53]. Most of these questions regard the stability of host-free LCC, and they are readily circumvented when the LCC are hosted by carbon nanotubes (CNT), when they are decorated with terminal groups such as the tris(3,5-di-*t*-butylphenyl)methyl, or when they are in colloidal environments [32,34,36-38,43,45,49-55]. Recently, single-wall (SW), double-wall (DW), and multiwall (MW) CNT have been used and considered ideal environments for fabricating stable LCC with up to 6000 carbon atoms [34,36-38,43,45,49-55]. Due to their 1D character, LCC are very simple structures, presenting rather simple electronic and phonon structures that are dependent on N [31,32]. When they are host-free, their phonon structures present longitudinal and transversal modes but their encapsulation by CNT inhibits transversal modes [29-32,55-59]. This inhibition seems to be confirmed in a recent work by Moura et al. [60], in which a novel Raman active longitudinal mode was observed for LCC, but no active transversal modes were observed despite theoretical predictions that suggested their existence.

The literature reports that LCC phonons possess long mean free paths (≈ 0.5 – $2.5 \mu\text{m}$) and lifetimes (≈ 30 – 110 ps), which tend to make ph–ph interactions inefficient [31,32]. These mean free paths and lifetimes are considerably larger when compared with other carbon materials [13,31-33]. In addition, several other works [29-32,55-59] have pointed out that CNT provide conditions that are sufficient to stabilize the LCC and inhibit transversal vibrations, while keeping CNT and LCC properties disentangled. In fact, this remains true even when the LCC@CNT systems are submitted to high pressures [29,55-57]. The literature has also shown that many chain-like quasi-1D materials constituted of C_{60} bulky-balls submitted to various conditions of pressure and temperature remain harmonic with properties that are independent of the properties from their hosting CNT [61-65]. This suggests that 1D-like materials also present inefficient ph–ph interactions, and that mutual interactions between chain-like structures and their hosts are second-order effects [61-65]. Sulfur chains inside SWCNT have also demonstrated enhanced field-emission properties and outstanding gas-sensing properties [66,67], which once again corroborate the idea that the hosting CNT primarily serve as a stabi-

lizing environment rather than one that alters the properties of the materials.

In this context, Costa and collaborators [30], demonstrated that LCC encapsulated by both multi-wall (LCC@MWCNT) and double-wall (LCC@DWCNT) carbon nanotubes are materials whose thermal properties can be described by the Debye model [30]. The reason is that the responses of materials to changing temperatures usually come from two contributions: (1) the lattice thermal expansion (LTE), associated with e–ph interactions; and (2) anharmonic effects, associated with ph–ph interactions. As is widely known, the Debye model does not consider ph–ph interactions but do describe quite well contributions associated with LTE [1,30]. Their work [30] was the first in the literature to provide experimental values for LCC internal energy per N (u), specific heat (c_v), coefficient of linear thermal expansion (α), thermal strain (ϵ_T), and Grüneisen parameter at constant pressure (γ_p).

The present work is intended to explore some important points that were left open by Costa and collaborators [30]. These points mainly regard the influence of the number of CNT walls in the LCC thermodynamic parameters as well as the responses of the system when measured isolated (or in very small bundles) and when measured in bundles. In addition, the current work extends the study to even lower temperatures (i.e., 13 K) when compared with previous data, whose minimum temperature stood around 70 K. Our analysis follows the same protocols described in reference [30]: the temperature evolution of the longitudinal optical phonon (so-called C-band), which is Raman active with frequencies (ω_{LCC}) around 1850 cm^{-1} , is thoroughly tracked and ω_{LCC} is used to indirectly access important thermodynamic parameters associated with LCC. Note that the

C-band is a spectroscopic signature widely used to identify distinct LCC since ω_{LCC} is dependent on the number of carbon atoms forming the chain (i.e., ω_{LCC} is size/length dependent).

Results and Discussion

The LCC encapsulated by multiwalled CNT (LCC@MWCNT) were synthesized using arc discharge [53]. The purity of MWCNT regarding nanoparticles is $\approx 80\%$ with average diameters of 10.4 nm (average length of 2.3 mm). The LCC@CNT filling ratio is $\approx 80\%$ [53]. The samples were dispersed in acetone and sonicated for 2 h and then drop casted onto a Si wafer of $\approx 1\text{ cm}^2$ area. Raman spectra were acquired with a $20\times$ objective lens in a backscattering geometry using Jobin Yvon Horiba T64000 spectrometer (1800 lines/mm grating). Samples were resonantly excited with 514.5 nm (2.41 eV) and 568.2 nm (2.18 eV) (Coherent Innova 70C Ar and Kr ion lasers).

The Raman spectra of LCC@MWCNT were acquired at low temperatures ranging from 13 to 293 K. Figure 1 shows representative spectra at 13 K and at 293 K, where temperature-dependent frequency shifts are clearly observed for both G- and C-bands (G-band comes from MWCNT). The LCC C-band was fitted with four Lorentzian curves for the spectra collected with the 514.5 nm (2.41 eV) excitation source, and two Lorentzian curves for the spectra collected with the 568.2 nm (2.18 eV) excitation source. Each Lorentzian represents a distinct LCC. Representative Raman spectra and respective fittings are shown in Supporting Information File 1, Figure S1. In Figure 2, N was estimated considering that ω_{LCC} is proportional to N^{-1} [29,51]. It is important to recall that the association between N and ω_{LCC} is only a reliable approximation. This number N is used here to correlate the length of LCC with the thermodynamic variables studied, and such approximation does not impair the analysis

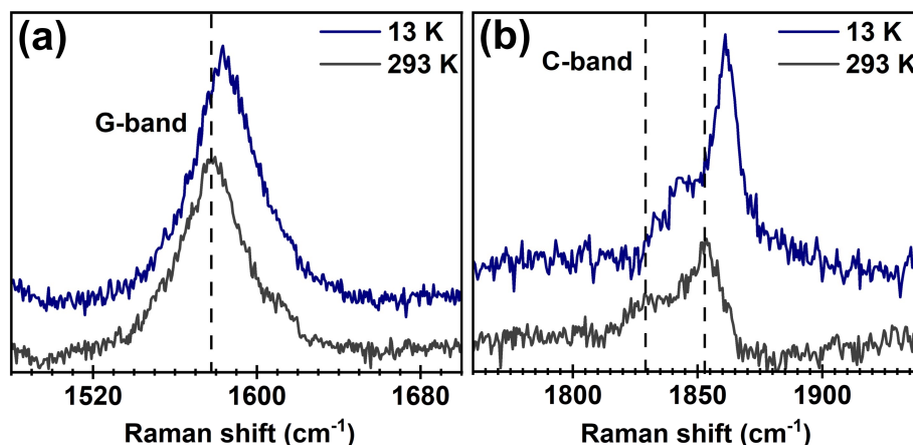


Figure 1: (a) and (b) show representative Raman spectra acquired from LCC@MWCNT at 13 K (solid navy blue curves) and 193 K (solid dark gray curves). The G-bands in (a) are associated with MWCNT, while the C-bands in (b) are associated with LCC. The vertical dashed lines are guides for the eyes. Supporting Information File 1, Figure S1 brings additional representative spectra.

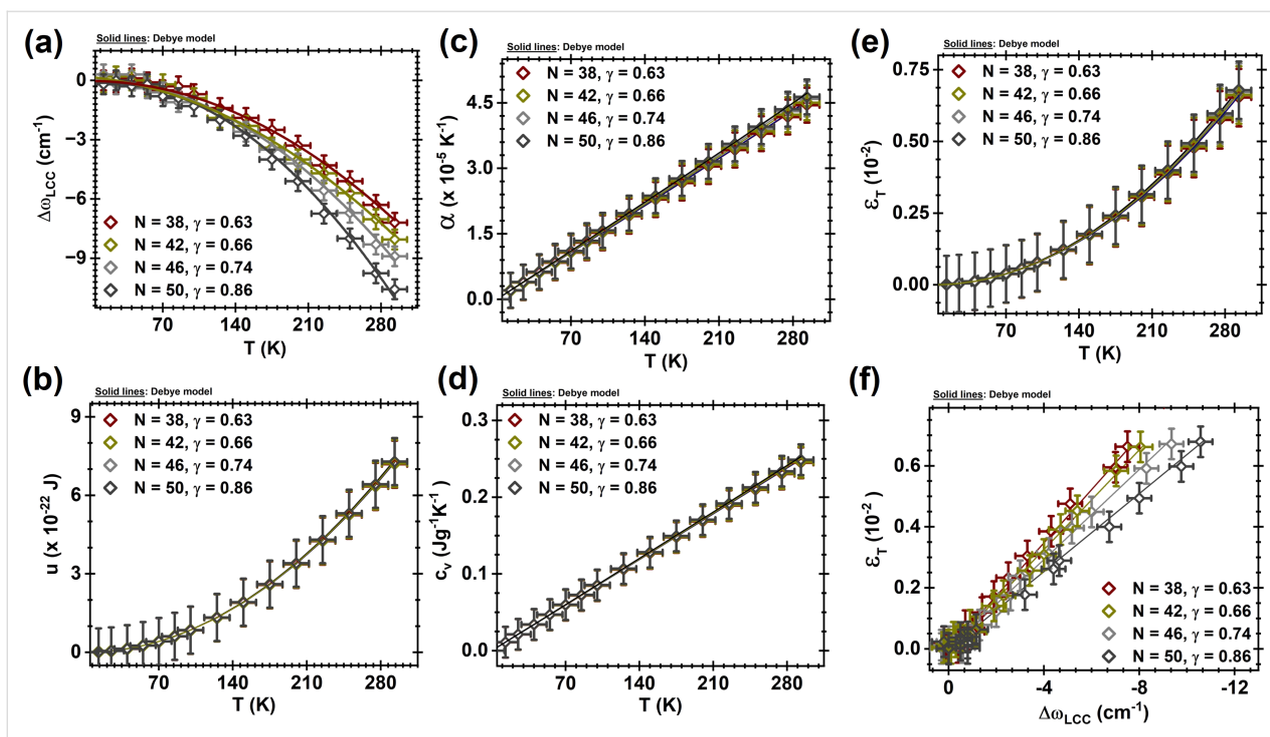


Figure 2: (a) Experimental $\Delta\omega_{\text{LCC}}(T)$ evolution with T ; (b) the energy per N , $u(T)$, presents a quadratic, universal, and unified behavior with T ; (c) $\alpha(T)$ shows a linear universal behavior with T ; (d) the heat capacity per N , $c_v(T)$, presents a linear, universal and unified behavior with T ; (e) A T^2 universal dependence is observed for the thermal strain $\varepsilon_T(T)$; (f) Every LCC presents a distinct linear dependence of ε_T with $\Delta\omega_{\text{LCC}}(T)$. Note that four representative LCC are shown here. The full set of LCC is presented in Figure S2 in Supporting Information File 1.

and conclusions of this paper. Finally, for the sake of clarity, Figure 2 shows results for representative LCC, while the full set of LCC is shown in Supporting Information File 1, Figure S2.

In total, four LCC with $N = 38, 42, 46,$ and 50 ($\gamma_P = 0.63, 0.66, 0.74,$ and 0.86) were identified using 514.5 nm, while two LCC with $N = 40$ and 50 ($\gamma_P = 0.67$ and 0.81 , respectively) were identified using 568.2 nm. In agreement with the literature [29,30], the use of different laser lines does not influence the response of the LCC@MWCNT to different T , but it might excite LCC with distinct N (LCC have their bandgap proportional to N^{-1} ; the smaller the chain, the larger the bandgap). Figure 2 corroborates this claim: ω_{LCC} for similar LCC possess similar dependence on T . As previously discussed, ω_{LCC} associated with each identified LCC is used as a probe to obtain the following thermodynamic properties as a function of T (T ranging from 13 to 293 K): $u(T)$, $\alpha(T)$, $c_v(T)$, $\varepsilon_T(T)$, and $\gamma_P(T)$.

The equations that correlate ω_{LCC} with these thermodynamic parameters are reported by Costa et al. [30] and reproduced in Supporting Information File 1 for reference (Equations S1–S6). Figure 2 shows a clear dependence of these properties with N and T , in accordance with previous work [30]. Costa and collaborators [30] studied these nanostructures under temperatures ranging from 70 to 293 K, but this work extends their results to

temperatures as low as 13 K. Here, we measured isolated LCC@MWCNT under 568.2 nm (2.18 eV) excitation, while those acquired under 514.5 nm (2.41 eV) excitation was in very small bundles. For reference, Costa et al. [30] measured small bundles in their work. The plots shown in Figure 2 provide an answer to one of the questions we sought to address in this work: do small bundles or isolated environments influence the vibrational and thermodynamic properties measured? Figure 2 suggests that the answer is no. In fact, it is evident that the data obtained in both scenarios are similar within the error margin expected in these experiments. In addition, the Raman spectra as well as the independent evolution as a function of temperature of the Raman bands from CNT and LCC (see Figure 1 and Figure S1 in Supporting Information File 1) suggest that the interaction between distinct LCC, and LCC and CNT are not strong enough to affect their electronic and phonon structures. Therefore, in agreement with the literature [29,30], mutual interactions between LCC and CNT are second-order effects [29,30]. Moreover, if ph–ph interactions can be neglected, the ω_{LCC} variation with $T(\Delta\omega_{\text{LCC}})$ should be well described by:

$$\Delta\omega_{\text{LCC}}(T) = \omega_{\text{LCC}}^0 \left(e^{-\gamma_P \int d\varepsilon} - 1 \right), \quad (1)$$

where ω_{LCC}^0 is the C-band frequency at $T = 0$ K, $d\epsilon = \alpha(T)dT$ is the thermal strain between T and $T + dT$, and γ_{P} is the T -independent Grüneisen parameter at constant P . Equation 1 in turn is expected to follow the empirical relation:

$$\Delta\omega_{\text{LCC}} = \Delta\omega_{\text{LCC}}(T) - \omega_{\text{LCC}}^0 = -\left(\frac{d^2\omega_{\text{LCC}}}{dT^2}\right)T^2, \quad (2)$$

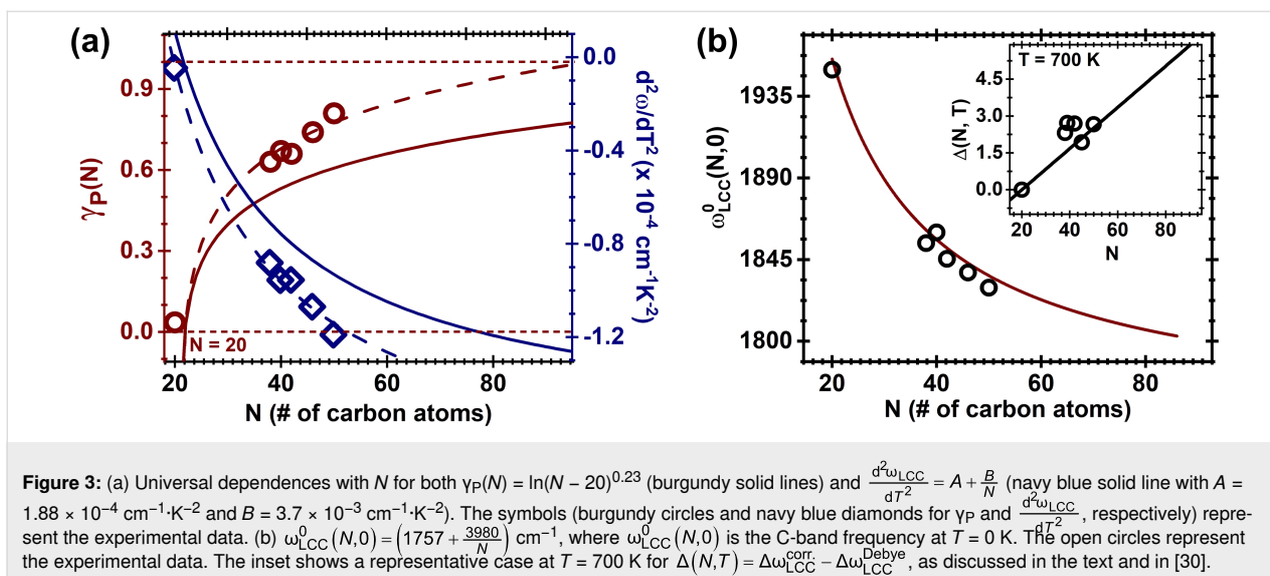
where the second derivative $\frac{d^2\omega_{\text{LCC}}}{dT^2}$ magnitude is N -dependent (see Figure 3). Figure 2a confirms that this is the case: the solid lines are fitting results using Equation 2. We are then well positioned to proceed with our analysis using the Debye model, whose associated equations (see Supporting Information File 1) depend on ω_{LCC} , $\frac{d\omega_{\text{LCC}}}{dT}$, and $\frac{d^2\omega_{\text{LCC}}}{dT^2}$ [30].

In one hand, Figure 2b shows that $u(T)$ displays a T^2 behavior; the values of $u(T)$ at 13 K, 70 K, and 293 K are 1.42×10^{-24} J, 4.16×10^{-23} J, and 7.27×10^{-22} J, respectively. On the other hand, $\alpha(T)$ and $c_{\text{v}}(T)$ (Figure 2c and 2d, respectively) vary linearly with T and do not show relevant dependence on N . Note that $\alpha(T)$ ranges from 4.40×10^{-5} K $^{-1}$ (293 K) to 1.98×10^{-6} K $^{-1}$ (13 K) with $\frac{d\alpha(T)}{dT} = 1.54 \times 10^{-7}$ K $^{-2}$, and $c_{\text{v}}(T)$ ranges from 0.25 J \cdot g $^{-1}$ \cdot K $^{-1}$ (293 K) to 0.01 J \cdot g $^{-1}$ \cdot K $^{-1}$ (13 K) with $\frac{dc_{\text{v}}(T)}{dT} = 8.84 \times 10^{-4}$ J \cdot g $^{-1}$ \cdot K $^{-2}$; both are in good agreement with the literature [30,37,68]. The heat extracted from the LCC leads their shrinkage, generating an internal pressure that is associated with the thermal strain (ϵ_{T}), which similarly with $u(T)$, displays a T^2 behavior (Figure 2e). This means that strains at 13 K (1.29×10^{-7} , this work) are very similar to those at 70 K (3.74×10^{-6}), while $\epsilon_{\text{T}} = 0.01 \times 10^{-2}$ at ambient conditions. As discussed in the literature [29,30,51], in addition to depending on the temperature, ω_{LCC} is dependent on the chain length (i.e., N -dependent) as well (see Figure 3b). This is an expected behavior since the size of the chain affects the bond length alternation (BLA) strength of the polyynes. Figure 2a shows the evolution of ω_{LCC} with T , and again, it is noteworthy that no matter the length of the chain, there is a convergence of the data as the temperature decreases. Consequently, ϵ_{T} as a function $\Delta\omega_{\text{LCC}}$ (see Figure 2f) also converges when $\omega_{\text{LCC}} \rightarrow \omega_{\text{LCC}}^0$. Note that in both cases, once again, the experimental data follow the Debye model up to 300 K. According to the data and the predictions above, $\Delta\omega_{\text{LCC}} \rightarrow 0$ and $\epsilon_{\text{T}} \rightarrow 0$ when $T \rightarrow 0$.

At this point, we are in good position to use Equation 1 to obtain γ_{P} associated with each measured chain, which in turn will deliver the γ_{P} dependence with N (Figure 3a). Note that once $\alpha(T)$ is known, γ_{P} becomes the only adjustable parameter in the equation. As anticipated earlier in the text, the found

values for γ_{P} are in accordance with those reported in previous works [29,30], endorsing inefficient phonon–phonon coupling in this system. The open symbols shown in Figure 3a display $\frac{d^2\omega_{\text{LCC}}}{dT^2}$ and γ_{P} as a function of N ; both of them displaying an universal dependence on N given, respectively, by $\gamma_{\text{P}}(N) = \ln(N - 20)^{0.23}$ and $\frac{d^2\omega_{\text{LCC}}}{dT^2} = A + \frac{B}{N}$, where $A = 1.88 \times 10^{-4}$ cm $^{-1}$ \cdot K $^{-2}$ and $B = 3.7 \times 10^{-3}$ cm $^{-1}$ \cdot K $^{-2}$ (dashed lines in Figure 3a). The solid lines in Figure 3a represent $\gamma_{\text{P}}(N)$ and $\frac{d^2\omega_{\text{LCC}}}{dT^2}(N)$ found by Costa et al. [30] for temperatures as low as 70 K. It is noticeable that there is a slight discrepancy between the data found in this work and those published by Costa and collaborators [30], which is explained as follows: as discussed, γ_{P} and $\frac{d^2\omega_{\text{LCC}}}{dT^2}$ are calculated from the fitting of the experimental data from Figure 1a, and they are, therefore, heavily dependent on $\omega_{\text{LCC}}(T, N)$ as a function of T , whose overall behavior is dependent on the range of temperatures used in the experiment. In the work by Costa et al. [30], the T range considered is limited to $70 < T < 300$ K, while this paper extends it to $13 < T < 300$ K. This newer range provides a more accurate prediction of ω_{LCC}^0 (i.e., $\omega_{\text{LCC}}(T)$ for $T = 0$ K), and therefore, the extra data we bring in this paper provides a better low-temperature convergent behavior of $\omega_{\text{LCC}}(N, T)$ and a better estimate for both $\frac{d\omega_{\text{LCC}}}{dT}$ and $\frac{d^2\omega_{\text{LCC}}}{dT^2}$, when compared with reference [30]. Since γ_{P} depends on $\Delta\omega_{\text{LCC}}(T)$, it is important to remind that, although the Debye model is understood to be valid even for temperatures beyond 300 K, the values we provide here are accurate up to 300 K only; these values of γ_{P} and $\frac{d^2\omega_{\text{LCC}}}{dT^2}$ might always be updated with a broader range of temperatures.

Finally, Kastner et al. [51] have predicted ω_{LCC}^0 to follow $\omega_{\text{LCC}}^0(N, 0) = 1757 + 3890/N$ (in cm $^{-1}$), which is plotted in the solid line of Figure 3b. The open circles represent our ω_{LCC}^0 values extrapolated from the experimental data, which is in good agreement with Kastner and collaborators. Also, the Debye model formalism works very well for temperatures as high as 300 K and corrections are predicted for $300 < T < 700$ K [30]. These corrections, however, continue to be small, as seen in the inset of Figure 3b, which plots the difference $\Delta(N, 700) = \omega_{\text{LCC}}^{\text{corr.}}(N, 700) - \omega_{\text{LCC}}^{\text{Debye}}(N, 700)$ for the representative case at 700 K. It is worth reminding that, as seen in Equations S1 to S6 in Supporting Information File 1, the superscript “Debye” stands for the predicted values without any corrections involving $\frac{d\omega_{\text{LCC}}}{dT}$ and $\frac{d^2\omega_{\text{LCC}}}{dT^2}$, while the superscript “corr.” stands for values that are corrected by such derivatives. The solid line represents the same correction as predicted by Costa and collaborators [30]. The dispersion of the data (open circles) with relation to the black solid curve has the same origins as those associated with the values of γ_{P} and $\frac{d^2\omega_{\text{LCC}}}{dT^2}$.



Conclusion

In summary, this paper investigates the thermodynamic properties of isolated and small bundles of LCC@MWCNT via Raman spectroscopy by tracking the C-band frequencies ω_{LCC} of LCC in the range of temperatures of $13 < T < 300 \text{ K}$. This range of temperatures provides more accurate values of $\frac{d\omega_{LCC}}{dT}$ and $\frac{d^2\omega_{LCC}}{dT^2}$, enhancing the reliability of the thermodynamic observables ($u(T)$, $\alpha(T)$, $c_v(T)$, $\varepsilon_T(T)$, and $\gamma_P(T)$). In addition, the data presented here further confirms that LCC may be well modelled using the Debye formalism even at ambient conditions. The thermodynamic observables indeed follow N -dependent universal laws with T . In this semiempirical model, the calculation of γ_P depends on the range of temperatures measured. The broader range of temperatures that this work considered allowed the authors to bring newly updated values of γ_P . These values might, however, undergo further corrections when more experimental data is available for temperatures beyond 300 K . This work also confirms that equivalent thermodynamic properties are observed for small bundles and isolates LCC@MWCNT.

Supporting Information

Supporting Information File 1

Additional figures and calculations.

[<https://www.beilstein-journals.org/bjnano/content/supplementary/2190-4286-16-125-S1.pdf>]

Funding

This material is based upon work supported by the National Science Foundation under Grant No. [1848418].

YAK acknowledges financial support from a National Research Foundation of Korea (NRF) grant funded by the Korea government (MSIP) (No. 2017M3A7B4014045). ARP and PTA would like to thank UFC CAPES-PrInt research proposal for the financial support in the scopes of 01/2018 and 41/2017 public calls from UFC and CAPES, respectively. ARP would like also to thank CNPq financial support in the scope of 04/2021 public call (grant No. 314084/2021-5).

Author Contributions

Alexandre Rocha Paschoal: conceptualization; data curation; formal analysis; funding acquisition; investigation; methodology; project administration; resources; supervision; validation; visualization; writing – original draft; writing – review & editing. Thiago Alves de Moura: investigation. Juan S. Rodríguez-Hernández: investigation. Carlos William de Araujo Paschoal: investigation; resources. Yoong Ahm Kim: resources. Morinobu Endo: resources. Paulo T. Araujo: conceptualization; data curation; formal analysis; funding acquisition; investigation; methodology; project administration; resources; supervision; validation; visualization; writing – original draft; writing – review & editing.

ORCID® iDs

Alexandre Rocha Paschoal - <https://orcid.org/0000-0002-7564-0884>

Thiago Alves de Moura - <https://orcid.org/0000-0002-7021-523X>

Juan S. Rodríguez-Hernández - <https://orcid.org/0000-0002-5137-4227>

Yoong Ahm Kim - <https://orcid.org/0000-0003-4074-7515>

Data Availability Statement

Additional research data generated and analyzed during this study is not shared.

References

- Araujo, P. T. *Phys. Rev. B* **2018**, *97*, 205441. doi:10.1103/physrevb.97.205441
- Stroscio, M. A.; Dutta, M. *Phonons in Nanostructures*; Cambridge University Press: New York, NY, USA, 2001. doi:10.1017/cbo9780511534898
- Klemens, P. G. *Phys. Rev.* **1966**, *148*, 845–848. doi:10.1103/physrev.148.845
- Castro Neto, A. H.; Guinea, F.; Peres, N. M. R.; Novoselov, K. S.; Geim, A. K. *Rev. Mod. Phys.* **2009**, *81*, 109–162. doi:10.1103/revmodphys.81.109
- Avouris, P.; Chen, Z.; Perebeinos, V. *Nat. Nanotechnol.* **2007**, *2*, 605–615. doi:10.1038/nnano.2007.300
- Balkanski, M.; Wallis, R. F.; Haro, E. *Phys. Rev. B* **1983**, *28*, 1928–1934. doi:10.1103/physrevb.28.1928
- Gao, K.; Dai, R.; Zhang, Z.; Ding, Z. *J. Phys.: Condens. Matter* **2007**, *19*, 486210. doi:10.1088/0953-8984/19/48/486210
- Viljas, J. K.; Heikkilä, T. T. *Phys. Rev. B* **2010**, *81*, 245404. doi:10.1103/physrevb.81.245404
- Shaina, P. R.; George, L.; Yadav, V.; Jaiswal, M. *J. Phys.: Condens. Matter* **2016**, *28*, 085301. doi:10.1088/0953-8984/28/8/085301
- Kong, B. D.; Paul, S.; Nardelli, M. B.; Kim, K. W. *Phys. Rev. B* **2009**, *80*, 033406. doi:10.1103/physrevb.80.033406
- Piscanec, S.; Lazzeri, M.; Mauri, F.; Ferrari, A. C.; Robertson, J. *Phys. Rev. Lett.* **2004**, *93*, 185503. doi:10.1103/physrevlett.93.185503
- Kang, K.; Abdula, D.; Cahill, D. G.; Shim, M. *Phys. Rev. B* **2010**, *81*, 165405. doi:10.1103/physrevb.81.165405
- Sendova, M.; Datas, L.; Flahaut, E. *J. Appl. Phys.* **2009**, *105*, 094312. doi:10.1063/1.3122301
- Malard, L. M.; Elias, D. C.; Alves, E. S.; Pimenta, M. A. *Phys. Rev. Lett.* **2008**, *101*, 257401. doi:10.1103/physrevlett.101.257401
- Araujo, P. T.; Mafra, D. L.; Sato, K.; Saito, R.; Kong, J.; Dresselhaus, M. S. *Phys. Rev. Lett.* **2012**, *109*, 046801. doi:10.1103/physrevlett.109.046801
- Tang, H.; Herman, I. P. *Phys. Rev. B* **1991**, *43*, 2299–2304. doi:10.1103/physrevb.43.2299
- Gao, B.; Hartland, G.; Fang, T.; Kelly, M.; Jena, D.; Xing, H. (Grace); Huang, L. *Nano Lett.* **2011**, *11*, 3184–3189. doi:10.1021/nl201397a
- Yoon, D.; Son, Y.-W.; Cheong, H. *Nano Lett.* **2011**, *11*, 3227–3231. doi:10.1021/nl201488g
- Calizo, I.; Balandin, A. A.; Bao, W.; Miao, F.; Lau, C. N. *Nano Lett.* **2007**, *7*, 2645–2649. doi:10.1021/nl071033g
- Magnin, Y.; Förster, G. D.; Rabilloud, F.; Calvo, F.; Zappelli, A.; Bichara, C. *J. Phys.: Condens. Matter* **2014**, *26*, 185401. doi:10.1088/0953-8984/26/18/185401
- Lee, J.-U.; Yoon, D.; Kim, H.; Lee, S. W.; Cheong, H. *Phys. Rev. B* **2011**, *83*, 081419. doi:10.1103/physrevb.83.081419
- Chen, S.; Moore, A. L.; Cai, W.; Suk, J. W.; An, J.; Mishra, C.; Amos, C.; Magnuson, C. W.; Kang, J.; Shi, L.; Ruoff, R. S. *ACS Nano* **2011**, *5*, 321–328. doi:10.1021/nn102915x
- Balandin, A. A. *Nat. Mater.* **2011**, *10*, 569–581. doi:10.1038/nmat3064
- Li, W. S.; Shen, Z. X.; Feng, Z. C.; Chua, S. J. *J. Appl. Phys.* **2000**, *87*, 3332–3337. doi:10.1063/1.372344
- Tse, W.-K.; Das Sarma, S. *Phys. Rev. B* **2009**, *79*, 235406. doi:10.1103/physrevb.79.235406
- Lindsay, L.; Broido, D. A.; Mingo, N. *Phys. Rev. B* **2011**, *83*, 235428. doi:10.1103/physrevb.83.235428
- Huang, L.; Hartland, G. V.; Chu, L.-Q.; Luxmi; Feenstra, R. M.; Lian, C.; Tahy, K.; Xing, H. *Nano Lett.* **2010**, *10*, 1308–1313. doi:10.1021/nl904106t
- Bonini, N.; Lazzeri, M.; Marzari, N.; Mauri, F. *Phys. Rev. Lett.* **2007**, *99*, 176802. doi:10.1103/physrevlett.99.176802
- Sharma, K.; Costa, N. L.; Kim, Y. A.; Muramatsu, H.; Neto, N. M. B.; Martins, L. G.; Kong, J.; Paschoal, A. R.; Araujo, P. T. *Phys. Rev. Lett.* **2020**, *125*, 105501. doi:10.1103/physrevlett.125.105501
- Costa, N. L.; Sharma, K.; Kim, Y. A.; Choi, G. B.; Endo, M.; Barbosa Neto, N. M.; Paschoal, A. R.; Araujo, P. T. *Phys. Rev. Lett.* **2021**, *126*, 125901. doi:10.1103/physrevlett.126.125901
- Wang, M.; Lin, S. *Sci. Rep.* **2016**, *5*, 18122. doi:10.1038/srep18122
- Kutrovskaya, S.; Osipov, A.; Baryshev, S.; Zasedatelev, A.; Samyshkin, V.; Demirchyan, S.; Pulci, O.; Grassano, D.; Gontrani, L.; Hartmann, R. R.; Portnoi, M. E.; Kucherik, A.; Lagoudakis, P. G.; Kavokin, A. *Nano Lett.* **2020**, *20*, 6502–6509. doi:10.1021/acs.nanolett.0c02244
- Swintek, N. Z.; Muralidharan, K.; Deymier, P. A. *J. Vib. Acoust.* **2013**, *135*, 041016. doi:10.1115/1.4023824
- Shi, L.; Rohringer, P.; Suenaga, K.; Niimi, Y.; Kotakoski, J.; Meyer, J. C.; Peterlik, H.; Wanko, M.; Cahangirov, S.; Rubio, A.; Lapin, Z. J.; Novotny, L.; Ayala, P.; Pichler, T. *Nat. Mater.* **2016**, *15*, 634–639. doi:10.1038/nmat4617
- Hirsch, A. *Nat. Mater.* **2010**, *9*, 868–871. doi:10.1038/nmat2885
- Zhao, X.; Ando, Y.; Liu, Y.; Jinno, M.; Suzuki, T. *Phys. Rev. Lett.* **2003**, *90*, 187401. doi:10.1103/physrevlett.90.187401
- Zhang, Y.; Su, Y.; Wang, L.; Kong, E. S.-W.; Chen, X.; Zhang, Y. *Nanoscale Res. Lett.* **2011**, *6*, 577. doi:10.1186/1556-276x-6-577
- Zhao, C.; Kitaura, R.; Hara, H.; Irlé, S.; Shinohara, H. *J. Phys. Chem. C* **2011**, *115*, 13166–13170. doi:10.1021/jp201647m
- Smith, P. P. K.; Buseck, P. R. *Science* **1982**, *216*, 984–986. doi:10.1126/science.216.4549.984
- Nair, A. K.; Cranford, S. W.; Buehler, M. J. *EPL* **2011**, *95*, 16002. doi:10.1209/0295-5075/95/16002
- Candiottio, G.; Silva, F. R.; Costa, D. G.; Capaz, R. B. *Phys. Rev. B* **2024**, *109*, 045405. doi:10.1103/physrevb.109.045405
- Lechner, J. M. A.; Marabotti, P.; Shi, L.; Pichler, T.; Casari, C. S.; Heeg, S. *Nat. Commun.* **2025**, *16*, 4360. doi:10.1038/s41467-025-59555-y
- Wang, Z.; Ke, X.; Zhu, Z.; Zhang, F.; Ruan, M.; Yang, J. *Phys. Rev. B* **2000**, *61*, R2472–R2474. doi:10.1103/physrevb.61.r2472
- Moura, L. G.; Malard, L. M.; Carneiro, M. A.; Venezuela, P.; Capaz, R. B.; Nishide, D.; Achiba, Y.; Shinohara, H.; Pimenta, M. A. *Phys. Rev. B* **2009**, *80*, 161401. doi:10.1103/physrevb.80.161401
- Shi, L.; Rohringer, P.; Wanko, M.; Rubio, A.; Waßerroth, S.; Reich, S.; Cambré, S.; Wenseleers, W.; Ayala, P.; Pichler, T. *Phys. Rev. Mater.* **2017**, *1*, 075601. doi:10.1103/physrevmaterials.1.075601
- Cretu, O.; Botello-Mendez, A. R.; Janowska, I.; Pham-Huu, C.; Charlier, J.-C.; Banhart, F. *Nano Lett.* **2013**, *13*, 3487–3493. doi:10.1021/nl4018918
- Lang, N. D.; Avouris, P. *Phys. Rev. Lett.* **1998**, *81*, 3515–3518. doi:10.1103/physrevlett.81.3515
- Lagow, R. J.; Kampa, J. J.; Wei, H.-C.; Battle, S. L.; Genge, J. W.; Laude, D. A.; Harper, C. J.; Bau, R.; Stevens, R. C.; Haw, J. F.; Munson, E. *Science* **1995**, *267*, 362–367. doi:10.1126/science.267.5196.362
- Nishide, D.; Dohi, H.; Wakabayashi, T.; Nishibori, E.; Aoyagi, S.; Ishida, M.; Kikuchi, S.; Kitaura, R.; Sugai, T.; Sakata, M.; Shinohara, H. *Chem. Phys. Lett.* **2006**, *428*, 356–360. doi:10.1016/j.cplett.2006.07.016

50. Cataldo, F. *Polyynes: Synthesis, Properties, and Applications*; CRC Press: Boca Raton, FL, USA, 2005. doi:10.1201/9781420027587.ch18
51. Kastner, J.; Kuzmany, H.; Kavan, L.; Dousek, F. P.; Kürti, J. *Macromolecules* **1995**, *28*, 344–353. doi:10.1021/ma00105a048
52. Diederich, F.; Kivala, M. *Adv. Mater. (Weinheim, Ger.)* **2010**, *22*, 803–812. doi:10.1002/adma.200902623
53. Chalifoux, W. A.; Tykwinski, R. R. *Nat. Chem.* **2010**, *2*, 967–971. doi:10.1038/nchem.828
54. Kang, C.-S.; Fujisawa, K.; Ko, Y.-I.; Muramatsu, H.; Hayashi, T.; Endo, M.; Kim, H. J.; Lim, D.; Kim, J. H.; Jung, Y. C.; Terrones, M.; Dresselhaus, M. S.; Kim, Y. A. *Carbon* **2016**, *107*, 217–224. doi:10.1016/j.carbon.2016.05.069
55. Andrade, N. F.; Vasconcelos, T. L.; Gouvea, C. P.; Archanjo, B. S.; Achete, C. A.; Kim, Y. A.; Endo, M.; Fantini, C.; Dresselhaus, M. S.; Souza Filho, A. G. *Carbon* **2015**, *90*, 172–180. doi:10.1016/j.carbon.2015.04.001
56. Andrade, N. F.; Aguiar, A. L.; Kim, Y. A.; Endo, M.; Freire, P. T. C.; Brunetto, G.; Galvão, D. S.; Dresselhaus, M. S.; Souza Filho, A. G. *J. Phys. Chem. C* **2015**, *119*, 10669–10676. doi:10.1021/acs.jpcc.5b00902
57. Neves, W. Q.; Alencar, R. S.; Ferreira, R. S.; Torres-Dias, A. C.; Andrade, N. F.; San-Miguel, A.; Kim, Y. A.; Endo, M.; Kim, D. W.; Muramatsu, H.; Aguiar, A. L.; Souza Filho, A. G. *Carbon* **2018**, *133*, 446–456. doi:10.1016/j.carbon.2018.01.084
58. Shi, L.; Sheng, L.; Yu, L.; An, K.; Ando, Y.; Zhao, X. *Nano Res.* **2011**, *4*, 759–766. doi:10.1007/s12274-011-0132-y
59. Endo, M.; Kim, Y. A.; Hayashi, T.; Muramatsu, H.; Terrones, M.; Saito, R.; Villalpando-Paez, F.; Chou, S. G.; Dresselhaus, M. S. *Small* **2006**, *2*, 1031–1036. doi:10.1002/sml.200600087
60. Moura, T. A.; Neves, W. Q.; Alencar, R. S.; Kim, Y. A.; Endo, M.; Vasconcelos, T. L.; Costa, D. G.; Candioto, G.; Capaz, R. B.; Araujo, P. T.; Souza Filho, A. G.; Paschoal, A. R. *Carbon* **2023**, *212*, 118123. doi:10.1016/j.carbon.2023.118123
61. Chorro, M.; Rols, S.; Cambedouzou, J.; Alvarez, L.; Almairac, R.; Sauvajol, J.-L.; Hodeau, J.-L.; Marques, L.; Mezouar, M.; Kataura, H. *Phys. Rev. B* **2006**, *74*, 205425. doi:10.1103/physrevb.74.205425
62. Rols, S.; Cambedouzou, J.; Chorro, M.; Schober, H.; Agafonov, V.; Launois, P.; Davydov, V.; Rakhmanina, A. V.; Kataura, H.; Sauvajol, J.-L. *Phys. Rev. Lett.* **2008**, *101*, 065507. doi:10.1103/physrevlett.101.065507
63. Cambedouzou, J.; Rols, S.; Almairac, R.; Sauvajol, J.-L.; Kataura, H.; Schober, H. *Phys. Rev. B* **2005**, *71*, 041403. doi:10.1103/physrevb.71.041403
64. Bousige, C.; Rols, S.; Ollivier, J.; Schober, H.; Fouquet, P.; Simeoni, G. G.; Agafonov, V.; Davydov, V.; Niimi, Y.; Suenaga, K.; Kataura, H.; Launois, P. *Phys. Rev. B* **2013**, *87*, 195438. doi:10.1103/physrevb.87.195438
65. Yang, J.; Lee, J.; Lee, J.; Yi, W. *Diamond Relat. Mater.* **2020**, *101*, 107554. doi:10.1016/j.diamond.2019.107554
66. Yang, J.; Lee, J.; Lee, J.; Yi, W. *Diamond Relat. Mater.* **2019**, *97*, 107474. doi:10.1016/j.diamond.2019.107474
67. Fujimori, T.; Morelos-Gómez, A.; Zhu, Z.; Muramatsu, H.; Futamura, R.; Urita, K.; Terrones, M.; Hayashi, T.; Endo, M.; Young Hong, S.; Chul Choi, Y.; Tománek, D.; Kaneko, K. *Nat. Commun.* **2013**, *4*, 2162. doi:10.1038/ncomms3162
68. Wong, C. H.; Buntov, E. A.; Rychkov, V. N.; Guseva, M. B.; Zatsopin, A. F. *Carbon* **2017**, *114*, 106–110. doi:10.1016/j.carbon.2016.12.009

License and Terms

This is an open access article licensed under the terms of the Beilstein-Institut Open Access License Agreement (<https://www.beilstein-journals.org/bjnano/terms>), which is identical to the Creative Commons Attribution 4.0 International License (<https://creativecommons.org/licenses/by/4.0>). The reuse of material under this license requires that the author(s), source and license are credited. Third-party material in this article could be subject to other licenses (typically indicated in the credit line), and in this case, users are required to obtain permission from the license holder to reuse the material.

The definitive version of this article is the electronic one which can be found at: <https://doi.org/10.3762/bjnano.16.125>