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Thermal effects on the characteristic Raman spectrum of molybdenum disulfide (MoS2) of varying thicknesses

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In this letter, thermal effects on the Raman spectra of molybdenum disulfide with thicknesses ranging from bulk to monolayer were evaluated. We quantitatively determined the laser-induced heating effects on the peak position and the line-width of the Raman spectrum. We found considerable thickness-dependent red-shifts as well as line-width changes for both $E_{2g}^1$ and $A_{1g}$ vibrating modes as laser power was increased. Our results enrich the knowledge of phononic behaviors of this material and demonstrate the important effects of the anharmonic terms in the lattice potential energy. © 2012 American Institute of Physics. [doi:10.1063/1.3673907]

Highly crystalline molybdenum disulfide (MoS2) films are composed of atomic layers of tightly bonded atoms brought together by weak van der Waals interlayer forces. These materials are structurally similar to graphene and hexagonal boron nitride (h-BN). Recent reports demonstrate strong photoluminescence emergence and anomalous lattice vibrations in single- and few-layered MoS2 films.\textsuperscript{1,2} This exemplifies the evolution of the physical and structural properties in MoS2, due to the transition from a three- to two-dimensional configuration. The thickness-dependency of the peaks can be explained by a combination of a classical model for the coupled harmonic oscillators\textsuperscript{3} and the anomalous changes in the dielectric screening properties of MoS2.\textsuperscript{4} Thickness-dependent changes are also reflected in the electronic properties of these materials. The indirect band gap of bulk MoS2 films with a magnitude of $\approx 1.2$ eV transforms gradually to a direct band gap of $\approx 1.8$ eV in single-layer samples.\textsuperscript{5,6} This is in contrast to pristine graphene with a band gap of $\approx 0$ eV and few-layered h-BN with a band gap of $\approx 5.5$ eV.\textsuperscript{7,8} The existence of an intrinsic band gap in these layered materials implies possible applications in electronics, optics, and semiconductor technologies as promising complements to graphene and h-BN.\textsuperscript{5–8}

To realize such technological possibilities, thermal characterization of these materials is vital. Raman spectroscopy has been widely applied to characterize the structural and physical properties of atomic layered materials, such as graphene and h-BN.\textsuperscript{9,10} However, characteristic Raman spectra of materials are known to be influenced by a number of extrinsic interactions, including effects caused by localized laser-induced heating, substrate interactions, strain, and charge transfer.\textsuperscript{11–17} Understanding these changes may provide important information regarding the underlying physical properties of the interactions and the materials themselves. Therefore, a thorough analysis considering thermal effects is necessary before utilizing Raman spectroscopy as a reliable characterization tool.

In this letter, we monitor the changes in the Raman spectrum of MoS2 caused by varied local heating, induced by the use of different laser powers. We demonstrate that the laser-induced heating generates a red-shift in both $E_{2g}^1$ and $A_{1g}$ vibrating modes of MoS2 samples. Similarly, the changes in the line-width of the Raman peaks are examined. Such changes are representative of the anharmonic terms in the lattice potential energy.\textsuperscript{17} This anharmonicity results in a shift in the peak position and widening of the line-width corresponding to the damping of the vibration.\textsuperscript{18} Both quantities are dependent on the heating effect by means of the thermal population factors of the interacting phonons.\textsuperscript{19}

High-quality MoS2 thin-film samples were prepared using both mechanical\textsuperscript{20} and liquid exfoliation methods.\textsuperscript{21} Single- and few-layered MoS2 films were obtained by both methods, and the samples can be viewed using an optical microscope. Single-, few-, and multi-layered MoS2 films are seen in light purple (similar to the color of SiO2 substrate), dark purple, and blue, respectively. Figures 1(a) and 1(b) show thin MoS2 flakes, with thicknesses ranging from a single layer to more than twenty layers. A SEM image of a liquid-exfoliated, few-layer MoS2 sample is illustrated in Figure 1(c). These samples’ thicknesses were further determined by atomic force microscopy (AFM, Agilent PicoScan 5500), as shown in Figures 1(d) and 1(e). The inset in Figure 1(d) depicts cross-sections of sample regions of thicknesses $\approx 0.7$ nm and 1.5 nm, corresponding to single- and bi-layered MoS2. The inset in Figure 1(e) shows thicknesses of $\approx 3.5$ nm and 6.5 nm MoS2 films, referred to as few- and multi-layered MoS2. The thickness of a liquid-exfoliated MoS2 flake may be determined by high-resolution transmission electron microscope (HRTEM, JEOL-2100). Figure 1(f) demonstrates a three-layer MoS2 sample, with layer spacing $\approx 0.7$ nm, and a Mo-Mo distance of about 0.31 nm, noted by the red dots. An atomic resolution HRTEM image of MoS2—clearly showing the hexagonal lattice structure—is shown in Figure 1(g). Figure 1(h) is the HRTEM image of MoS2 flakes on carbon film and the reduced-fast Fourier transform (R-FFT) of these samples, demonstrating the diffraction pattern of a single-layered MoS2 sample.

As shown in Figure 2(a), two characteristic Raman active modes for mechanical- and liquid-exfoliated MoS2—$E_{2g}^1$ 383 cm$^{-1}$ and $A_{1g}$ 409 cm$^{-1}$—are present in the spectrum. These vibrational modes have been theoretically and experimentally investigated in bulk MoS2.\textsuperscript{22,23} The inset of Figure 2(a) demonstrates the Raman active planar vibrations $E_{1g}$ and $E_{2g}^1$, as well as $A_{1g}$, which associates with the vibration modes.
of sulfides in the out-of-plane direction. It is believed that $E_{1g}$ peak is not visible because of the limited rejection of the Rayleigh scattered radiation by the SiO$_2$ substrate. A laser with an excitation wavelength of 514.5 nm (Renishaw inVia) was used to examine the thickness-dependence of the MoS$_2$ Raman spectrum. Similar to previously reported results, the thickness-dependent changes in the Raman spectra of MoS$_2$ samples on SiO$_2$ substrates were observed.\textsuperscript{2} These results illustrate that the $E_{1g}^2$ and $A_{1g}$ peaks move closer together as the thickness of pristine MoS$_2$ decreases, i.e., $A_{1g}$ has a red-shift in the order of 5 cm$^{-1}$, while $E_{1g}^1$ has a blue-shift of approximately 2 cm$^{-1}$ from bulk- to single-layered MoS$_2$. A rate increase for these changes is observed from bi- to single-layered samples.

Thermal effects on the Raman spectrum of the MoS$_2$ samples on SiO$_2$ substrate were evaluated. It is known that laser power can be used to effectively control the heat-generating interactions between light and mater.\textsuperscript{24} To illustrate the thermal effects in the Raman spectra of MoS$_2$, samples of different thicknesses ranging from bulk to single atomic layer at high (20 mW), intermediate (8 mW), and low (2 mW) laser power levels were examined. The resulting difference in the peak position, obtained for the varied laser powers, clearly signifies a considerable thickness-dependent change in the laser-induced thermal effects. These differences are represented by an expected softening of bonds (Fig. 2(b)). In thick samples (number of layers $N > 6$ to bulk), negligible changes in both peak positions is seen (Fig. 2(c)). In thinner samples ($N < 6$), we observed a red-shift for both $A_{1g}$ and $E_{1g}^2$ modes. The rate of this shift for both modes of vibration is roughly equal in many- and few-layered samples. An increasing discrepancy between the magnitudes of the red-shifts for the two modes of vibration is observed as the number of layers is decreased. This is believed to be a reflection of the thickness-dependency of the Raman spectrum due to the thermal effects. A decrease in material thickness results in the stiffening of the $E_{1g}^1$ mode, which contributes to the gradual mitigation of the thermal softening rate.\textsuperscript{2,4} In contrast, the $A_{1g}$ mode softens with a decrease in thickness, resulting in enhanced thermal softening. In single-layered samples, the maximum difference between the responses of the two peaks is measured, and the change in the peak position of $A_{1g}$ mode is about 1 cm$^{-1}$ larger than $E_{1g}^1$. This can be explained by the observed increased stiffing of $E_{1g}^2$ and increased softening of $A_{1g}$ modes from bi- to single-layered MoS$_2$.

It has been shown that the evolution of line-width, resulting from changes in the thermal effects, can provide important information about phonon-phonon coupling, as well as the decay process and population number of phonons in the material.\textsuperscript{25} Here, we have measured the full width at half maximum (FWHM) of the peaks and have observed their change as laser power is increased (Figs. 3(a) and 3(b)). Results obtained from a low-power laser are in accordance with a previous study of the thickness-dependence of the line-width.\textsuperscript{2} We observed that enhanced thermal effects result in an amplified phonon coupling of both peaks, represented by an increased line-width as the laser power is increased. The reduced sensitivity to thickness of the $E_{1g}^2$ mode has been explained by the less-efficient inter-layer coupling of the in-plane phonons.\textsuperscript{2} However, this is subjected to change as the laser power is increased; thicker samples are more susceptible to such couplings (Fig. 3(a)). For the $A_{1g}$ mode at low laser powers, the maximum line-width

![HRTEM of a single-layer MoS$_2$ on SiO$_2$ substrate.](image)\textsuperscript{013106-2} Najmaei \textit{et al.}\textsuperscript{1} Appl. Phys. Lett. 100, 013106 (2012)
is observed for the bi-layered samples, and a gradual decrease in the line-width is observed as thickness increases. The rapid line-width, decreases from bi- to single-layered samples, has been explained by an intrinsic difference in their symmetry. The thermally induced rate of change in the line-width of the $A_{1g}$ mode is highest in single-layered samples. As a result, single-layered samples obtain the maximum line-width value at high laser powers (Fig. 3(b)). Thus, due to heating, $A_{1g}$ phonon coupling is most rapidly increased in single-layered samples.

In conclusion, our results suggest that the Raman spectrum of MoS$_2$ shows significant temperature dependency. We reveal that both characteristic peaks, $A_{1g}$ and $E_{1g}^1$, are sensitive to temperature changes, which are reflected in the peak position and line-width of the Raman spectra. This behavior is thickness-dependent, and a dramatic difference is
observed for single-layered, relative to bi-layered, samples. The findings from this work reveal the importance of considering thermal effects on characterization of these materials and enrich our knowledge of the phononic behavior of MoS$_2$ materials with varying thicknesses.

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