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| Author(s) | Luo, Zhiqiang; Cong, Chunxiao; Zhang, Jun; Xiong, Qihua; Yu, Ting |
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Zhiqiang Luo, Chunxiao Cong, Jun Zhang, Qihua Xiong, and Ting Yu

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Direct observation of inner and outer G' band double-resonance Raman scattering in free standing graphene

Zhiqiang Luo,¹ Chunxiao Cong,¹ Jun Zhang,¹ Qihua Xiong,¹ and Ting Yu¹,²,³,a)
¹Division of Physics and Applied Physics, School of Physical and Mathematical Sciences, Nanyang Technological University, Singapore, 637371
²Department of Physics, Faculty of Science, National University of Singapore, Singapore, 117542
³Graphene Research Centre, National University of Singapore, Singapore, 117542

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In contrary to the widely reported single and symmetric peak feature of G' band in Raman spectrum of graphene, we herein report the observation of splitting in G' band in free standing graphene. Our experimental findings provide a direct and strong support for the previous theoretical prediction that the coexistence of the outer and inner processes in the double-resonance Raman scattering would cause the splitting of G' mode. The investigation of the influence of trigonal warping effect on the spectral features of G' subbands further verified the theoretical interpretation established on the anisotropic electronic structure of graphene.

Raman scattering has been playing an important role in the study of structural and electronic properties of graphene,¹⁻⁷ which is a mono-layer of graphite showing many intriguing physical properties. Two dominant characteristic Raman features usually present in a Raman spectrum of graphene, the so called G band and G' (or 2D) band.¹ The G band originates from a single resonance process associated with doubly degenerate iTO and LO phonon modes at the Brillouin zone center (Γ point), while G' band is associated with two phonon intervalley double resonance (DR) scattering involving iTO phonon near K point.¹ It is widely accepted that the G' band is composed of a single, sharp, and symmetric Lorentzian peak for single layer graphene.²,⁶,⁷ However, the theoretical calculation of the G' bands of graphene suggested that the G' band of graphene should be in a split form, since both the outer process in KΓ direction and the inner process in KM direction should play significant contribution in DR Raman scattering.²,⁹ Direct observation of the splitting of the G' band is thus important for verifying these theoretical results and understanding of the detailed DR Raman scattering process in graphene. In this work, using free standing graphene samples, we observed the splitting of G' band and further investigated the influence of trigonal warping effect on these two G' subbands.

The free standing graphene samples were prepared by micromechanical exfoliation of highly ordered pyrolytic graphite onto a SiO₂/Si substrate pre-patterned with an array of trenches, which was fabricated by photolithography and reactive ion etching.¹⁰ The Raman spectra were recorded by Renishaw inVia Raman system with excitation lasers of 2.33 eV (532 nm) and 1.58 eV (785 nm), and Jonin-Yvon T64000 Raman system with excitation lasers of 3.49 eV (535 nm) and 1.96 eV (633 nm). The laser power on graphene sample is kept below 1 mW to avoid possible laser-induced heating.

Figure 1(a) shows an optical image of a typical free standing graphene sample used in this study. For all the free standing graphene samples, there is no noticeable strain in the suspended area as evidenced by the negligible shift of G band compared to those of the non-suspended area³,¹⁰ (see Raman spectra in Figure 1(b)). It is interesting to notice that, the G' band of the free standing graphene is obviously asymmetric, while the G' band of the graphene on SiO₂/Si substrate appears a broader symmetric peak. The Lorentzian peak fittings of the highly asymmetric G' peak recorded under the excitation of 2.33 eV is displayed in Figure 1(c).

In most of the previous reports, the observed G' band of graphene on a substrate can always be nicely fitted with one Lorentzian line.²,⁶,⁷ However, in the intervalley DR Raman scattering of the G' band, there are two possible scattering processes, so called outer process and inner process.⁸,¹¹ As illustrated in Figures 2(a) and 2(b), the outer process is associated with electron transition in KT direction, while the inner process is associated with electron transition in KM direction.⁸,¹² Due to the trigonal warping effect,¹³ the energy dispersion of electron in KT direction is steeper. The outer process involves the photo-excited electron with smaller momentum, and then phonon with lower frequency, while the inner process involves the photo-excited electron with larger momentum and phonon with higher frequency.⁸,¹² It would be appropriate to assign the G'₁ and G'₂ to the outer process and the inner process, respectively. On the basis of peak fitting, the intensity ratio of the inner to the outer processes was estimated of approximately 35% ~ 60%. The smaller weight of the inner process would come from the lower phonon density of states in the KT direction.¹²

a)Author to whom correspondence should be addressed. Electronic mail: Yuting@ntu.edu.sg.

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The trigonal warping effect in graphene is expected to show significant influence on the spectral features associated with the outer and inner process. Actually, there is a subtle change in the full width at half-maximum (FWHM) of $G_1^0$ and $G_2^0$ at different excitation energies. As shown in Figure 3(a), for the excitation of 1.58 eV, the FWHM of the $G_1^0$ and $G_2^0$ peaks are around 19 cm$^{-1}$. However, for the excitation of 2.33 eV, the FWHM of the $G_1^0$ and $G_2^0$ peaks are around 15.5 cm$^{-1}$ and 23 cm$^{-1}$, respectively (see Figure 1(c)). The deviation of the FWHM of the outer and inner process at higher excitation energy should be attributed to the enhanced trigonal warping effect at higher energy level (see illustration in Figures 2(c) and 2(d)).

For the photo-electrons at the same energy level, deviance of the change in electron momentum ($Dk$) during electron scattering is smaller in the flatter edge along the $\Gamma M$ direction, while it becomes larger in the sharper curvature in $KM$ direction. The smaller deviation of the change in electron momentum at $\Gamma K$ direction will result in smaller FWHM of the Raman peak corresponding to the outer process (momentum of the phonon $q = Dk$).

The trigonal warping effect causes an apparent polarization dependence of the inner to outer processes ratio (see Figure 3). Taking the $G'$ band excited by 1.58 eV laser as an example, when the excitation and detection polarizations are parallel ($\theta = 0^\circ$), the inner to outer processes ratio is about 35%. Whereas, when the excitation and detection...
polarizations are orthogonal \((\theta = 90^\circ)\), the inner to outer processes ratio is about 45\%. Obviously, as shown in Figures 3(c) and 3(d), this polarization dependence becomes more apparent in the UV region. For better understanding of this polarization dependent inner to outer processes ratio, polarized Raman spectra involving of G band and G’ band are shown in Figure 4. When \(\theta = 0^\circ\), the \(I_{G'}/I_G = 7.5\), and \(I_{G''}/I_G = 2.5\), respectively. When \(\theta = 90^\circ\), the \(I_{G'}/I_G = 3.3\), \(I_{G''}/I_G = 2.1\), and \(I_{G''}/I_G = 1.2\), respectively. It is well known that the G band intensity of graphene is isotropic; however, due to the inhomogeneous optical absorption and emission in DR Raman scattering process associated with the G’ band, the G’ band intensity of graphene is anisotropic: \(I_{G'}(\theta) = [2\cos^2(\theta) + 1]I_0/3\), where \(\theta\) is the angle between the polarization of the analyzer and the polarization of the incident laser, and the \(I_0\) is a maximum Raman intensity for the G’ band when \(\theta = 0^\circ\). The \(I_{G'}(\theta)/I_{G'}(\theta = 90^\circ) = 3.3/10 = 1/3\) in our measurement is in good agreement with the calculation result. The \(I_{G'}(\theta)\) is the intensity integrated over the whole Brillouin Zone, involving the contribution from both the outer and the inner processes. The polarization dependence of the intensity of the separated G’ subbands corresponding to the outer and inner processes should not simply follow the \(I_{G'}(\theta)\).

The absorption probability in the light absorption process as a function of angle around the K point shows significant difference for the outer and inner processes, which should result in different polarization dependence for G’ subbands. The electron states around KM direction with sharper curvature have higher absorption probability accompany with wider angle distribution. In the inner process, wider angle distribution of optical absorption at KM direction should cause smaller weight in the projection of the scattered light \((P_s)\) along the polarization direction of incident light \((P_L)\), and therefore the \((I_{G'}/I_G)_{\theta = 90^\circ}/(I_{G'}/I_G)_{\theta = 0^\circ} = 0.28\) and \((I_{G''}/I_G)_{\theta = 90^\circ}/(I_{G''}/I_G)_{\theta = 0^\circ} = 0.48\), which well fits the above theoretical picture. It is clear that the polarization dependence of the inner to outer processes ratio results from the trigonal warping effect.

The broader symmetric G’ band of graphene on SiO2/Si substrate should result from the merging of the two G’ subbands due to the increased Raman broadening parameter induced by substrate doping. It is worth to point out that
when the graphene on a substrate was subject to a sufficient strain, the splitting of G' band had also been observed, which would be another support of coexistence of both outer process and inner process. In this situation, the strain induced asymmetry of the Brillouin zone play important role in revealing of G' sub-bands with large FWHM, and thus the spectral features are quite sensitive to the direction of incident laser polarization with respect to the strain axis. However, there is no significant strain in the free standing graphene, and the observed G' band splitting is, therefore, a more direct evidence of the coexistence of outer and inner processes.

In summary, the two well fitted Lorentzian lines in our observed asymmetric G' band of the free standing graphene is a direct evidence of the coexistence of both outer process and inner process in the DR Raman scattering. The trigonal warping effect in electronic structure of graphene shows significant influence on the FWHM and polarization dependence of the subbands associated with the outer and inner processes, which well supports the theoretical interpretation established on the anisotropic electronic structure of graphene. This interesting observation experimentally reveals the detailed Raman scattering process corresponding to the G' band in graphene.

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