Drude conductivity of Dirac fermions in graphene

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Electrons moving in graphene behave as massless Dirac fermions, and they exhibit fascinating low-frequency electrical transport phenomena. Their dynamic response, however, is little known at frequencies above one terahertz (THz). Such knowledge is important not only for a deeper understanding of the Dirac electron quantum transport, but also for graphene applications in ultrahigh-speed THz electronics and infrared (IR) optoelectronics. In this paper, we report measurements of high-frequency conductivity of graphene from THz to mid-IR at different carrier concentrations. The conductivity exhibits Drude-like frequency dependence and increases dramatically at THz frequencies, but its absolute strength is lower than theoretical predictions. This anomalous reduction of free-electron oscillator strength is corroborated by corresponding changes in graphene interband transitions, as required by the sum rule.

I. INTRODUCTION

Graphene provides a unique material system to study Dirac fermion physics in two dimensions. Researchers have demonstrated in graphene exotic Dirac fermion phenomena ranging from anomalous quantum Hall effects1,2 to Klein tunneling3 in low-frequency (dc) electrical transport. They also observed an optical conductance defined by the fine-structure constant1,2 and gate-tunable infrared (IR) absorption in Dirac fermion interband transitions.4,5 Situated between low-frequency electrical transport and interband optical excitation is the spectral range dominated by free-carrier intraband transitions. Infrared spectroscopy study on free-carrier responses of epitaxial graphene layers has been performed by several groups and provides new insight into both Dirac fermion transport and interband transitions.4,5 This free-carrier dynamics response is also expected to play a key role in the future development of ultrahigh-speed electronics at terahertz (THz) frequencies and THz-to-mid-IR optoelectronic devices.6,7 However, despite its importance, an experimental study of free-carrier response of graphene in the THz-to-mid-IR spectral range is still lacking.

Quantum theories of electron dynamic response in graphene have been developed by several groups.12−17 They predict that, within the framework of Boltzmann transport theories, high-frequency conductivity of monolayer graphene has a Drude form, σ(ω) = [D/π(ω + iΓ)], where ω is the frequency and Γ the scattering rate. The prefactor D, known as the Drude weight, has the value D = (v_F^2 e^2/h)√π|n| when electron-electron interactions are neglected, with v_F = 1.1 × 10^6 m/s being the Fermi velocity and |n| the carrier density. This is distinctly different from classical materials where D = πne^2/m^*.

The predictions on graphene Dirac fermion dynamics have never been directly tested experimentally. In fact, it was suggested from measurements of interband transitions6 that D could deviate from the predicted value. There are also theoretical studies showing that electron-impurity interactions could result in carrier dynamics in graphene beyond the Boltzmann description18,19 and that inclusion of electron-electron interactions could dramatically reduce the Drude weight D.20

II. EXPERIMENTAL

We report spectroscopic measurements of graphene conductivity from terahertz to mid-IR for different electron and hole concentrations. We confirm that the conductivity from free-carrier response indeed has a Drude-like frequency dependence. In addition, we have been able to determine the Drude weight D and scattering rate Γ of doped graphene directly and separately. We observe an electron-hole asymmetry for D, and we find that values of D are lower than the theoretical prediction D = (v_F^2 e^2/h)√π|n|.12−14 This reduction of Drude weight (arising from free-carrier intraband transitions) is connected to corresponding changes in interband transitions, and we demonstrate that the sum rule requiring the integrated intraband (Drude) conductivity and interband conductivity to be a constant is well obeyed.

In our study, we used Fourier-transform infrared spectroscopy (FTIR) to measure the transmission spectra of graphene samples over the range from 30 to 6000 cm−1 and deduce the frequency-dependent conductivity from the spectra. Previously, infrared spectroscopy to probe a graphene monolayer was limited to >1000 cm−1 (wavelength < 10 μm) because of the limited size of exfoliated graphene.4,6,7,23 For spectroscopy at a longer wavelength up to ~300 μm, we need a large sample size. Here we used large-area graphene grown by chemical vapor deposition (CVD).

Following the procedure in Ref. 22, we grew graphene on copper films using CH_4 as the feed gas, which was then transferred with PMMA (Poly(methyl methacrylate)) support to a Si/SiO_2 wafer after wet-etching to remove the copper film by FeCl_3. After dissolving the PMMA in acetone solution,
measurements. The doped Si substrate (through stencil masks onto the graphene sample for electrical transitions up to 2
to a SiO2/Si substrate. (b) Graphene dc conductivity as a function of gate voltage. The conductivity minimum at Vg =
14 V defines the charge neutral point (CNP). (c) Gate-induced change of ir transmittance ΔT/T through graphene at Vg = −70 V (hole
compared to transmittance at the CNP. The spectrum shows an increase of free-carrier absorption at low wave numbers and a
reduction of interband absorption at higher wave numbers. (d) An illustration of intraband (i.e., free-carrier absorption, dashed arrow) and
interband (solid arrow) transitions in hole-doped graphene. Intraband absorption increases with carrier doping, while interband transitions up to 2EF become forbidden due to empty initial states, as observed in (c).

high-quality graphene of centimeter size on the Si/SiO2 wafer was obtained [Fig. 1(a)]. Raman spectra show that the
sample was indeed monolayer graphene. Subsequently, Au/Ti electrodes (thickness ~50 nm) were deposited in vacuum through stencil masks onto the graphene sample for electrical measurements. The doped Si substrate (p-type, resistivity ~10 Ω cm) underneath the 290 nm SiO2 served as a back gate, which allowed us to tune the charge-carrier density in the graphene sample. All optical and electrical measurements were performed in vacuum (~0.1 mTorr) at 100 K.

III. RESULTS AND DISCUSSION

Figure 1(b) shows the dc conductivity as a function of the gate voltage in a typical graphene sample. The minimum conduc-
tance occurs at 14 V, which defines the charge neutral point (CNP) of the sample. Carrier density in graphene is related to the
gate voltage by n = 7.5 × 10^{10}(V_{CNP} − V_g) cm^−2 V^−1, which where negative (positive) density corresponds to electron
(hole) doping. To double-check this relationship, carrier concentration is also measured by the infrared spectroscopy method developed from Ref. 6, in which we identify the onset energy of interband absorption at 2EF at a different gate voltage and then compute the concentration by $E_F = \hbar v_F \sqrt{\pi |n|}$. The carrier density obtained from two different methods shows that systematic error for the carrier concentration is less than 5%. The sample had an initial hole doping of 1.05 × 10^{12} cm^−2 at Vg = 0 and a hole mobility around 2700 cm^2/V s. As seen from Fig. 1(b), the dc conductivity for hole doping is reasonably linear with |n|, and there is a large asymmetry between electron and hole conductivity at the same |n|. In previous studies, conclusions have frequently been made on carrier scattering from such dc conductivity data based on $\sigma_{dc} = D/πΓ$ and the assumption that $D = (v_F e^2/h)\sqrt{\pi |n|}$ holds exactly as theory predicts. However, the validity of this approach has never been tested.

An independent determination of D and Γ can be achieved through ac conductivity measurements using IR spectroscopy. Figure 1(c) shows a difference IR absorption spectrum of hole-doped graphene ($V_g = −70$ V) in reference to absorption at the CNP ($V_g = 14$ V). This difference absorption spectrum has two characteristic features: a large absorption increase at lower wave numbers, which reaches over 15% at terahertz frequencies for a graphene monolayer, and an absorption reduction over a broad range of higher wave numbers. These features can be understood qualitatively from the gate-induced changes in intraband and interband electronic transitions in graphene due to carrier doping [Fig. 1(d)]. Free-carrier conductivity (from intraband transitions) increases dramatically with the hole doping. It peaks at zero frequency and gives rise to a substantial absorption increase at low wave numbers. At the same time, however, the interband transitions up to energy $2EF$ (arrows in Fig. 1(d)] become forbidden due to empty initial states, leading to a reduction of absorption in the broad spectral range below $2EF$ (dashed line in Fig. 1(e)).

The gate-induced change of ac conductivity (referred to as the CNP value), $\Delta\sigma = \sigma - \sigma_{CNP}$, can be obtained readily from the difference IR absorption spectra (see auxiliary material). Figures 2(a) and 2(b) show the real part of the ac conductivity change $\Delta\sigma^\prime$ in the low-wave-number range ($< 450$ cm^−1) for different electron and hole concentrations. In this spectral range, free-carrier response (i.e., an increase in conductivity with carrier density) dominates, providing direct information on Dirac fermion electrical transport. The complete set of $\Delta\sigma^\prime$ spectra in Figs. 2(a) and 2(b) can be fit (dashed line) by $\Delta\sigma^\prime(\omega) = i D/\pi(\omega + i\Gamma) − i D_{CNP}/\pi(\omega + i\Gamma_{CNP})$ using the Drude form for $\sigma$ and $\sigma_{CNP}$. A finite CNP response ($D_{CNP}$ and $\Gamma_{CNP}$) accounts for inhomogeneous electron and hole puddles present in graphene. In the fittings, we set $\Gamma_{CNP}$ equal to that when graphene is weakly doped. The value of $D_{CNP}$ is set so that $D_{CNP}/\pi\Gamma_{CNP}$ gives the dc conductance at the CNP. From the Drude fits, we obtain directly the scattering rate $\Gamma$ and Drude weight $D$ as a function of electron and hole concentrations, shown as symbols in Figs. 2(c) and 2(d), respectively.

From the values of D and $Γ$ extracted from the IR spectroscopy, we can obtain $σ_{dc}(= D/πΓ)$ of graphene at different carrier concentrations [red dots in Fig. 2(e)]. It agrees nicely with the directly measured dc conductivity in our sample for all gate voltages (black dots). This agreement provides a consistency check of the Drude description.

We now compare our experimentally determined $Γ$ and D to theory predictions. Figure 2(c) shows that $Γ$ is roughly a constant at different hole concentrations, while it is higher in value and increases with the carrier concentration for electron doping. This electron-hole concentration dependence
Gate-induced change of interband ac conductivity $\Delta \sigma'$ in hole- and electron-doped graphene, respectively, over 600 $< \omega < 6000$ cm$^{-1}$. The ac conductivity decreases due to empty initial states (hole doping) or filled final states (electron doping) for interband transitions below 2$E_F$. (The sharp features around 1200 cm$^{-1}$ are extrinsic and arise from SiO$_2$ phonon resonances of the substrate.) Observed interband conductivity decrease is appreciably less than the theoretically predicted value of $\pi e^2/2h$ (horizontal dashed line), and shows electron-hole asymmetry. (c) Upon charge-carrier doping, integrated conductivity change has an increase from intraband transitions (squares) that equals the decrease from interband transitions (circles). The two values agree quantitatively at all gate voltages, as required by the oscillator strength sum rule.

FIG. 3. (Color online) Interband transitions in graphene. (a,b) Gate-induced change of interband ac conductivity $\Delta \sigma'$ in hole- and electron-doped graphene, respectively, over 600 $< \omega < 6000$ cm$^{-1}$. The ac conductivity decreases due to empty initial states (hole doping) or filled final states (electron doping) for interband transitions below 2$E_F$. (The sharp features around 1200 cm$^{-1}$ are extrinsic and arise from SiO$_2$ phonon resonances of the substrate.) Observed interband conductivity decrease is appreciably less than the theoretically predicted value of $\pi e^2/2h$ (horizontal dashed line), and shows electron-hole asymmetry. (c) Upon charge-carrier doping, integrated conductivity change has an increase from intraband transitions (squares) that equals the decrease from interband transitions (circles). The two values agree quantitatively at all gate voltages, as required by the oscillator strength sum rule.

The general sum rule for oscillator strengths in solids states that the integrated conductivity over all frequencies is a constant, i.e., $\int_0^{\infty} \Delta \sigma' \omega d\omega = 0$. In the case of noninteracting free carriers described by prevailing graphene theories, the integrated intraband absorption increases by $\int \Delta \sigma'_\text{int} d\omega = D/2 = (v_F e^2/2h)\sqrt{\pi n}$ upon carrier doping. This increase is compensated exactly by an integrated interband absorption $\int \Delta \sigma'_\text{inter} d\omega = \sigma_0(2E_F/h)\sqrt{\pi n}$ with $E_F = h\nu\sqrt{\pi n}$, and $\sigma_0 = e^2/4h$. In our case, the sum rule is still valid, and the observed anomalous reduction of Drude weight in intraband transitions should be accompanied by a corresponding change in interband transitions at large wave numbers. Figures 3(a) and 3(b) show the optical conductivity difference (referred to as CNP) at different hole and electron concentrations in the spectral range 600–6000 cm$^{-1}$. A gate-induced decrease in optical conductivity is clearly present from zero to a cutoff frequency of $2E_F/h$ that increases with carrier doping, as the theory predicts. However, the observed interband optical conductivity decrease in doped graphene is appreciably less than $\sigma_0$ (dashed lines in Figs. 3(a) and 3(b)), and the deviation is larger for electron doping compared with hole doping. This reduction in gate-induced absorption decrease, as well as electron-hole asymmetry in interband transitions, matches well with the reduction in gate-induced absorption increase and electron-hole asymmetry in intraband Drude weight. To be more quantitative, we plot in Fig. 3(c) the absolute values of integrated conductivity change for interband transitions (black symbol) and intraband transitions (red symbol, which is proportional to the Drude weight change) at different carrier doping. The two agree almost perfectly, just as the sum rule requires. A reduction of gate-induced
interband absorption has also been observed previously for mechanically exfoliated graphene.\textsuperscript{6} Based on the sum rule that we established experimentally, it indicates that the anomalous Drude weight reduction is general for both exfoliated and CVD samples.

This reduction of Drude weight cannot be easily understood. One theoretical study showed that electron-electron interactions in graphene could lead to a reduced Drude weight.\textsuperscript{18} However, this theory predicted a much larger reduction of the Drude weight (over 80%) for graphene on SiO\(_2\) than what we have observed. It is also possible that a reduction of \(D\) is a consequence of defect-induced electron or hole localization, which decreases the effective “free” electron or hole concentration. In addition, the electron-impurity interactions may play a role and contribute to the observed electron-hole asymmetry.\textsuperscript{19} Further experiments with higher-quality graphene will be required to pin down the underlying mechanism for the Drude weight reduction.

IV. SUMMARY

Our infrared spectroscopy shows that the frequency-dependent response of a Dirac fermion in graphene can be described by the Drude form, which becomes very strong dependent on SiO\(_2\) than what we have observed. It is also possible that a reduction of \(D\) is a consequence of defect-induced electron or hole localization, which decreases the effective “free” electron or hole concentration. In addition, the electron-impurity interactions may play a role and contribute to the observed electron-hole asymmetry.\textsuperscript{19} Further experiments with higher-quality graphene will be required to pin down the underlying mechanism for the Drude weight reduction.

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APPENDIX: CALCULATING THE AC CONDUCTIVITY

The frequency-dependent conductivity change of graphene \(\Delta \sigma\) at different gate voltages is obtained from the measured transmission spectra using a perturbation treatment, because a monolayer graphene only absorbs a small fraction of the light. Within this approximation, the complex ac conductivity change is related to the difference transmission spectra by

\[
\frac{-\Delta \tilde{T}(\omega)}{T} = \frac{4\pi}{c} \text{Re}(\Delta \tilde{\sigma} \times L),
\]

where \(-\Delta \tilde{T}/T\) is the normalized transmission difference, \(L\) is the local field factor, and \(\Delta \tilde{\sigma}\) is the gate-induced conductivity change in graphene.

For suspended graphene, the local field factor is 1, and we obtain the well-known form of \((-\Delta T/T)(\omega) = (4\pi/c)\Delta \sigma\)\textsuperscript{18}. In our device, the graphene is sitting on a SiO\(_2\)/Si substrate and the local field factor \(L\) is uniquely defined by the refractive index of Si and SiO\(_2\), as well as the thickness of SiO\(_2\), and it can be determined accurately without any adjusting parameters. Specifically, \(L\) has the form

\[
L = 1 + r_{ag} + t_{ag}r_{gs}t_{ga} - e^{2ikn_d}r_{ag},
\]

where \(r_{ag}, r_{gs},\) and \(t_{ga}\) are the Fresnel reflection coefficients at air-silica, silica-silicon, and silica-air interferences, \(r_{ag}\) and \(t_{ga}\) are the corresponding transmission coefficients, \(n\) and \(d\) are the complex reflective index and thickness of the SiO\(_2\) layer, respectively, and \(k\) is the wave vector of the incident light.

The real and imaginary parts of ac conductivity, \(\sigma'\) and \(\sigma''\), are connected by the Kramers-Kronig (KK) relation

\[
\sigma''(\omega) = -\frac{2\omega}{\pi} \int_0^\infty \sigma'(\tilde{\omega})d\tilde{\omega} / \tilde{\omega}^2.
\]

Because the KK relation holds true for all the gate voltages, the gate-induced conductivity change \(\Delta \sigma = \Delta \sigma' + i \Delta \sigma''\) also satisfies the same KK relation. Making use of Eq. (A1) and the KK relation, we obtain both the real and imaginary parts of the gate-induced ac conductivity from infrared transmission spectra.

The gate-induced infrared absorption from the \(p\)-doped Si substrate is negligible. The real part of hole ac conductivity in silicon is described by \(\sigma' = (ne^2/m_p)[\Gamma^2 + \omega^2]\). For the same carrier density, it is more than one order of magnitude smaller than graphene conductivity across the experimental spectral range (30 < \(\omega < 6000\) cm\(^{-1}\)) due to the large hole effective mass (\(m_h = 0.36m_0\)) and small scattering rate (\(\Gamma \sim 4\) cm\(^{-1}\) at 100 K).\textsuperscript{31}

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