Ultrafast Dynamics of Massive Dirac Fermions in Bilayer Graphene

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Bilayer graphene is a highly promising material for electronic and optoelectronic applications since it is supporting massive Dirac fermions with a tunable band gap. However, no consistent picture of the gap’s effect on the optical and transport behavior has emerged so far, and it has been proposed that the insulating nature of the gap could be compromised by unavoidable structural defects, by topological in-gap states, or that the electronic structure could be altogether changed by many-body effects. Here, we directly follow the excited carriers in bilayer graphene on a femtosecond time scale, using ultrafast time- and angle-resolved photoemission. We find a behavior consistent with a single-particle band gap. Compared to monolayer graphene, the existence of this band gap leads to an increased carrier lifetime in the minimum of the lowest conduction band. This is in sharp contrast to the second substate of the conduction band, in which the excited electrons decay through fast, phonon-assisted interband transitions.

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The lack of a band gap is the most important obstacle to using graphene in electronic devices but this can be elegantly solved in bilayer graphene (BLG) when an asymmetry between the layers is induced by a transverse electric field [1–3]. The promising properties of the thereby induced massive Dirac particles have been intensively explored for the development of semiconducting devices with tunable band gaps [4–6] and for efficient photodetectors extending to the THz regime [7–10]. However, the quantitative transport properties of BLG are inconsistent with the simple scenario of a small band gap semiconductor [5] and different hypotheses for this have been given, including broken-symmetry ground states [11] or topological edge state effects [12]. Recently, a study of BLG by angle-resolved photoemission spectroscopy (ARPES) has revealed the presence of electronic states throughout the gap, arising from intrinsic AA-stacked domains [13], something that might be expected to short circuit the gap of BLG.

While static ARPES results provide crucial information about the spectral function of BLG, time-resolved spectroscopy is needed to complement the transport studies. Time- and angle-resolved photoemission (TR-ARPES) experiments near the Dirac cone have only recently become technically feasible [14,15] due to the high photon energies needed to reach the K point in the Brillouin zone. Here, we report TR-ARPES measurements carried out using a Ti:sapphire amplified laser system with a repetition rate of 1 kHz, carried out at the Artemis facility at the Central Laser Facility, Rutherford Appleton Laboratory [16]. This provided ultrafast infrared pulses with a wavelength of 785 nm, a full width at half maximum duration of 30 fs, and an energy per pulse of 12 mJ. A part of the laser energy was applied for high harmonic generation of extreme ultraviolet pulses in a pulsed jet of argon gas. A time-preserving monochromator was used to single out the 13th harmonic with a photon energy of 21 eV. The remaining laser energy was utilized to drive an optical parametric amplifier, which can provide tunable pump pulses. We used the second harmonic of the signal to produce 30 fs pulses at 1.55 eV. Both beams were polarized perpendicular to the scattering plane. The data were acquired by sweeping 45 time delay points. Such a cycle was repeated between 700 and 1000 times in order to obtain satisfactory statistics in the data, amounting to total acquisition times between 8 and 14 h per data set.

Samples of so-called quasifreestanding mono- and bilayer graphene were prepared ex situ by thermal decomposition of SiC substrates followed by hydrogen intercalation to decouple the graphene layers from the substrate [17]. Both samples are hole doped with carrier concentration on the order of 5 × 10¹² cm⁻², which places the
FIG. 1 (color online). Ultrafast ARPES measurements of bilayer graphene: (a) Normalized photoemission intensity integrated above the Fermi level. Vertical dashed lines mark the times of the dispersion snapshots in (g)–(j). (b)–(e) Diagrams of ultrafast processes and relaxation dynamcis involving optical pumping (straight arrows), electron scattering (curled arrows), optical (vertical blue wiggled arrows), and acoustic (green wiggled arrows) phonon scattering. Filled (open) circles signify electrons (holes). (f) Spectrum before the arrival of the pump pulse with the inset showing the acquisition direction for all data (arrows), and acoustic (green wiggled arrows) phonon scattering. Filled (open) circles signify electrons (holes). (g)–(j) ARPES difference spectra obtained by subtracting the spectrum in (f) from the spectrum taken at the given time. A pump laser fluence of $F = 1.0 \text{ mJ/cm}^2$ was used. The dashed parabolic bands are the result of a tight binding calculation and have been added as a guide to the eye.

Dirac point 240 meV above the Fermi level in the monolayer sample, while in the bilayer a substrate-induced charge asymmetry between the two graphene layers opens a band gap centered 180 meV above the Fermi level [3,13]. Both samples were transferred through air into the TR-ARPES ultrahigh vacuum end station, where they were cleaned by annealing to 550 K. The samples were held at 90 K throughout the experiment.

Figure 1(a) presents the integrated intensity of photoinduced excited states above the Fermi level during and immediately after the pumping phase. The pump signal induces direct transitions from the two parabolic valence bands to their conduction band counterparts. This pumping phase is characterized by a fast rising edge [see Fig. 1(a)]. The excited electrons quickly thermalize, consistent with rapid Auger-like processes [14,18,19] as depicted in Fig. 1(b). This leaves a dense transient population of hot electrons in both conduction bands as sketched in Fig. 1(c). The electrons then quickly cascade down to the region around the band gap, thereby depleting the conduction bands by emission of phonons [Fig. 1(d)]. This process continues [Fig. 1(e)] until all of the excited electrons have recombined with the remaining holes.

Snapshots of these events are presented in Figs. 1(g)–1(j), displaying the ARPES difference spectra between the equilibrium signal before the arrival of the pump [Fig. 1(f)] and the signal at the given time delay. The bare band dispersion lines obtained by a tight binding calculation (see Refs. [1,20]) are plotted on top of the difference spectra to match the intensity with the band structure. The two bilayer bands are clearly discerned in the raw data as well as in the valence band part of the difference spectra. Note that we do not observe discrete spikes in the intensity distribution at the energies corresponding to the direct transition, not even the midpoint of the rising edge at $t = 0 \text{ fs}$ [see Fig. 1(g)], but rather an even intensity distribution along the bands. This implies that the initial thermalization of the hot electrons by electron-electron scattering occurs on time scales less than 40 fs, too fast to be resolved here. Once the peak distribution is reached at $t = 40 \text{ fs}$ [see Fig. 1(h)], the phonon scattering processes lead to a fast decrease of intensity as observed in Figs. 1(i)–1(j).

Within the considered time window, the described behavior is qualitatively identical for the laser pump fluence of $F = 1.0 \text{ mJ/cm}^2$ used for acquiring the data in Fig. 1 and for another data set with $F = 3.5 \text{ mJ/cm}^2$ [20]. However, with the higher fluence, a larger density of photoexcited electrons is created, which allows us to study a larger population of hot electrons in the two conduction bands. A quantitative study of this is introduced in Fig. 2.
FIG. 2 (color online). Disentangling the dynamics in the bilayer bands: (a),(e) Photoemission intensity integrated between the dashed lines in (b) and (f) as a function of binding energy. Markers correspond to data points at the given time delay, and lines are integrated intensity from simulated photoemission data with the given temperatures. (b),(f) Experimental photoemission data at the peak excitation signal at time $t = 40$ fs. (c),(g) Simulated data at the given temperature taking the experimental resolution into account. (d),(h) Simulated data without resolution broadening. The vertical dashed lines in (a) and (e) and the horizontal dashed lines in (d) and (h) mark the onset energy of the two conduction bands. The data in (a)–(d) correspond to a fluence of 1.0 mJ/cm$^2$ while the data in (e)–(h) correspond to 3.5 mJ/cm$^2$.

Here, we compare the time-dependent experimental data with simulations where we populate the conduction bands thermally by multiplying a model spectral function with a single high temperature Fermi-Dirac (FD) function [20].

We extract the combined statistical distribution of the carriers in the four bands at a given time delay by integrating the photoemission intensity in between the dashed lines in Figs. 2(b) and 2(f) [21]. Interestingly, the integrated distribution curves (IDCs) of the transient spectra presented in Figs. 2(a) and 2(e) do not appear to be merely described by a featureless hot electron FD tail, but a clear shoulder can be discerned in the raw data. The extent of this feature is a function of time and of fluence. As it is not possible to extract the temperature from the data by a fit to a simple FD function, we match the observed photoemission intensity to a simulation, consisting of the calculated spectral function that is multiplied by a hot FD function and broadened by the experimental resolution. Note that the intensity variation between the bands and within the bands originate from the photoemission matrix elements that are accounted for in the simulated spectral functions. These matrix elements are assumed to be energy and band dependent but independent of the fluence or time [20].

The smearing of the separation between the conduction bands in the simulation is consistent with the measured data, and can be attributed to the combined effect of resolution and matrix elements. We find that the electrons are heated from the initial temperature of 90 K to reach temperatures of 2 900 K and 4 200 K for low and high fluence, respectively.

Remarkably, the complex shape of the IDCs from the peak signal at $t = 40$ fs and onwards in time can be described by this simple simulation. The observed shoulder in the IDCs always appears at the same energy which coincides with the onset of the upper conduction band, as seen by comparing the IDCs to the nonbroadened BLG spectral function via the dashed lines in Figs. 2(a), 2(d), 2(e), and 2(h). The shoulder can, therefore, be ascribed to the accumulation of electrons in the upper conduction band.

The fact that all the IDCs can be fitted with this simple model also suggests that the excited carriers in both conduction bands are in a quasiequilibrium state within our time resolution.

The model presented in Fig. 2 integrates over all the subbands in the system and can, thus, neither be used to study possible differences in the subbands, nor describe lifetime effects that do not follow a FD distribution. A more detailed analysis and a direct comparison to the situation in monolayer graphene (MLG) is, therefore, shown in Fig. 3. For both materials, the Dirac point imposes a bottleneck for relaxation processes, but the key consequence of a true band gap in BLG would be to impose a serious constriction for the decay of hot electrons, e.g., by scattering with low-energy acoustic phonons. To avoid any complications from high fluence effects [20], we consider both MLG and BLG at a low fluence of 0.6 mJ/cm$^2$. The equilibrium spectra for BLG and MLG at this fluence are shown in Figs. 3(a) and 3(f), and the difference spectra from $t = 40$ fs to $t = 2000$ fs are shown in Figs. 3(b)–3(d) and 3(g)–3(i), respectively.

The time scales of the relaxation processes are directly extracted by analyzing the intensity binned in small boxes following the dispersion, as shown in the difference spectra in Fig. 3. Contrary to the simple model in Fig. 2, this provides access to the spectrally resolved dynamics at specified binding energies [22], capturing dynamics that would otherwise be integrated out. The time-dependent intensity for all boxes is presented in Figs. 3(e), 3(j), and 3(k). It can be described by a simple exponential decay, apart from box 1, the box of the conduction band minimum in both samples (Note that the location of box 1 has been defined such that it is at equal distance from the center of the gap and the Dirac point in BLG and MLG, respectively). We shall discuss the intensity in this spectral region later.

The electron population in the upper conduction band in BLG is tracked by boxes 5 and 6 in Fig. 3(e), and we find a fast decay with a time constant of 300 and 200 fs, respectively. For box 6, this is expected because the electrons can cascade down to the minimum of the upper...
There is still intensity from hot electrons in box 1 while in already be seen in the raw data in Figs. The pronounced difference between the two materials can of 10 000 fs merely sets a lower limit for the long decay time. Possible within our experimental time window and the value slow that a precise determination of the decay constant is not (400 fs) for BLG (MLG) is followed by a slow decay with decay where an initial fast decay with a decay time of 650 fs Both data sets can be well-described by a double-exponential band, pronounced differences for BLG and MLG are found. For this spectral region, the very bottom of the conduction band, pronounced differences for BLG and MLG are also similar.

FIG. 3 (color online). Comparing the relaxation dynamics of massive and massless Dirac Fermions: (a) Spectrum before the arrival of the pump pulse and (b)–(d) difference spectra at selected time delays for BLG. (f)–(i) Corresponding data for MLG. Dispersion lines from tight binding models have been added as guide to the eye. (e),(j),(k) Normalized photoemission intensity binned over the boxes in (b)–(d) for (e) BLG and (g)–(i) for (j) MLG. The time traces in (e) and (j) have been labeled with a number according to the box numbers, and the given time constants are results of single exponential fits to the corresponding trace. The error bar on these fits amounts to ±20 fs. (k) Integrated intensity in box 1 for both BLG and MLG. Lines correspond to double-exponential function fits with the given time constants. The fluence is 0.6 ml/cm² for all data presented here. (l),(m) Sketch of the relaxation dynamics involving hot electrons (red spheres), holes (yellow spheres), and phonon emission (wiggled arrows) around the Dirac point for (l) massive Dirac Fermions and (m) massless Dirac Fermions.

Conduction band. For box 5, the fast decay can only be explained by phonon-induced interband scattering to the lower conduction band. A similarly fast decay is found for corresponding boxes in the lower conduction band (boxes 3 and 4), suggesting that the interband scattering is sufficiently effective to avoid any electron accumulation near the upper conduction band edge. Indeed, this is also responsible for the fact that the simple FD fit works in Fig. 2. The decay times for the corresponding energies in MLG are also similar.

The situation is more complicated for box 1 in both materials where the intensity decay cannot be fit to a single exponential. The time-dependent intensity of box 1 is shown in Fig. 3(k) up to a measured delay time of 4 ps. For this spectral region, the very bottom of the conduction band, pronounced differences for BLG and MLG are found. Both data sets can be well-described by a double-exponential decay where an initial fast decay with a decay time of 650 fs (400 fs) for BLG (MLG) is followed by a slow decay with 10 000 fs (3 500 fs). Indeed, the slow decay for BLG is so slow that a precise determination of the decay constant is not possible within our experimental time window and the value of 10 000 fs merely sets a lower limit for the long decay time. The pronounced difference between the two materials can already be seen in the raw data in Figs. 3(d) and 3(i). In BLG, there is still intensity from hot electrons in box 1 while in MLG the whole spectrum is fully relaxed.

The true difference in the hot electron carrier dynamics thus occurs mainly at the Dirac point—an experimental verification that is only possible using this direct momentum-resolved technique. This result is consistent with the naive expectation of a substantially longer decay time in the presence of a gap. Without a gap, hot electrons can cascade down continuously via acoustic phonon scattering, whereas the presence of the gap requires relaxations by optical phonons, giving rise to a bottleneck for the decay [see sketch in Figs. 3(l) and 3(m)]. The efficiency of a relaxation across the gap might be further restricted by the role of the pseudospin in BLG [1].

In view of the disputed character of the electronic structure of BLG and the role of possible in-gap states, we should also point out that the spectral function fits and the data analysis in Fig. 3 are perfectly consistent with the expected single-particle tight-binding band structure. The possibility of in-gap states still exists, especially that of gap-crossing states in AA-stacked domains, since these were identified for the same kind of BLG samples used here [13]. In order to obtain a quantitative estimate of the impact these defects have on the lifetime of the carriers, a calculation of the carrier lifetime for the perfect crystalline system would be required. Nevertheless, the results shown here reveal that, even in the presence of such gap states, the band gap leads to a markedly enhanced lifetime over MLG.

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