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We show with angle-resolved photoemission spectroscopy that a new energy band appears in the electronic structure of electron-doped hydrogenated monolayer graphene (H-graphene). Its occupation can be controlled with the hydrogen amount and allows for tuning of graphene’s doping level. Our calculations of the electronic structure of H-graphene suggest that this state is largely composed of hydrogen 1s orbitals and remains extended for low H coverages despite the random chemisorption of H. Further evidence for the existence of a hydrogen state is provided by x-ray absorption studies of undoped H-graphene which are clearly showing the emergence of an additional state in the vicinity of the $\pi^*$ resonance.

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I. INTRODUCTION

Doping is at the heart of modern semiconductor technology because it allows for control of the carrier density and is therefore the basis for all circuit elements. If the concentration of the dopant increases beyond a critical value, the physical properties of a doped semiconductor are described by impurity band formation. For example in the case of B-doped diamond an insulator-to-metal transition and superconductivity is observed beyond a critical B concentration. From angle-resolved photoemission spectroscopy (ARPES) and x-ray absorption spectroscopy of B-doped diamond, it was concluded that the impurity band derived from B orbitals is above $E_F$. Covalent doping is also successfully applied to sp2 bonded carbon materials. It has been shown that hydrogen readily forms a covalent bond with graphene. The bonding environment of C-H is usually well defined and H/C ratios equal to $\sim$25% have been realized. It is therefore tempting to ask whether impurity band formation happens in hydrogenated graphene (H-graphene), in close analogy to B-doped diamond. Theory suggests H-graphene to have a substantial band gap of 1 eV accompanied by a dispersionless and spin-polarized midgap state. Transport experiments have shown that the exposure of graphene to atomic hydrogen turns graphene into an insulator with a temperature dependence of the conductivity that points toward a variable range hopping mechanism. For electron-doped ($n$-doped) graphene on SiC, small H amounts yield a shrinkage of the Fermi surface, and a metal-to-insulator transition (MIT) has been observed with a combined transport and ARPES study. For pristine quasi-free-standing graphene, the opening of a band gap has been reported upon hydrogenation. However, the formation of a hydrogen impurity band has not yet been observed experimentally and therefore the question of its existence and electronic properties remains an open topic.

In this work we present the first experimental evidence of a hydrogen-derived midgap state in H-graphene using ARPES and near-edge x-ray absorption fine structure (NEXAFS) measurements. The latter method provides access to unoccupied electronic states above the Fermi level by measuring the x-ray absorption which is caused by transitions from the C 1s core level to the $\pi^*$ and $\sigma^*$ energy bands. For undoped H-graphene the impurity band is located within the emerging gap at the Fermi level and is therefore hardly accessible with ARPES since this method probes only the occupied electronic states.
of the band structure. Using potassium intercalated $n$-doped graphene, the hydrogen-derived state becomes available for an occupation with electrons and acts as an acceptor level for $\pi^*$ electrons which can be directly observed in ARPES.

Density functional theory (DFT) calculations suggest that the midgap state is largely derived from H 1$s$ orbitals. Calculations of the typical and average density of states (DOS) using the kernel polynomial method (KPM) with a tight-binding (TB) band structure calculation\textsuperscript{11,16} indicate that the new state does not localize easily despite the randomness in the H chemisorption sites.

II. EXPERIMENTAL METHODS

Pristine monolayer graphene samples were prepared in situ under ultrahigh vacuum conditions by chemical vapor deposition on Ni(111) thin films epitaxial grown on W(110).\textsuperscript{17} Then one monolayer (ML) of Au was deposited on graphene and intercalated into the graphene/Ni interface by annealing:\textsuperscript{7,18,19} This procedure liberates graphene from the strong interaction to the Ni substrate, rendering it quasi–free standing. Functionalization was performed by exposing pristine graphene to potassium atoms which were evaporated from commercial SAES metal dispensers while the graphene sample was kept at 25 K during the deposition. Potassium intercalation into the graphene/Au interface was performed at room temperature (RT). Hydrogenation was carried out at $\sim 1 \times 10^{-9}$ mbar for times between 1 and 100 s. The atomic hydrogen beam was produced by cracking H$_2$ at 3000 K in a tungsten capillary. ARPES measurements were carried out at the BaDElPh beamline of the Elettra synchrotron in Trieste (Italy).\textsuperscript{20} The spectra were acquired at photon energies of 26 eV and 40 eV with the sample at 25 K and a base pressure better than $8 \times 10^{-11}$ mbar. The angular resolution was 0.15° and the energy resolution was set to 15 meV. All ARPES measurements were taken along the $\Gamma K M$ directions. Furthermore, we performed NEXAFS measurements at the carbon K edge using synchrotron radiation. These studies were carried out at BESSY II using the HESGM beamline equipped with a home-built channeltron detector. The NEXAFS spectra were measured in the partial electron yield mode. We have applied a negative retarding potential of $U = -150$ V.

III. RESULTS AND DISCUSSION

We discuss the electronic properties of hydrogenated graphene intercalated with Au using NEXAFS. As this method probes the unoccupied density of states by inducing dipole transitions from the C 1$s$ core levels to the $\pi^*$ energy band, it is a versatile technique to investigate the changes in the electronic structure that appear upon hydrogenation. In Figs. 1(a) and 1(b) we show the NEXAFS spectra for pristine and H-graphene (H/C $\sim$ 15%),\textsuperscript{7} respectively. Both sets of curves depict a similar behavior with $\phi$, the angle of light incidence: For grazing incidence ($\phi = 20^\circ$) we have a maximum photoabsorption referring to the $\pi^*$ resonance and for normal incidence ($\phi = 90^\circ$) we have a maximum absorption identified as the $\sigma^*$ resonance. This is in agreement with the NEXAFS experiments performed on graphite and previous measurements on graphene.\textsuperscript{7,21,22} A closer look at the $\pi^*$ resonance is shown in Fig. 1(c). Two observations can be made from the comparison of pristine and H-graphene: (1) the $\pi^*$ resonance gets weaker in intensity upon hydrogenation and (2) the high-resolution spectra show a low-energy shoulder appearing upon hydrogenation for grazing incidence. Observation 1 can be well understood since hydrogen adsorption effectively removes $\pi$ bonds ($s p^2$ C-H bond). Concerning observation 2, the feature appears between 284.3 eV and 285.3 eV, as indicated by the arrows in Fig. 1(c). In the following we argue that this extra feature can also be attributed to a midgap state that lies in between the $\pi$ and $\pi^*$ bands. Such a feature can contribute to a downshift of the $\pi^*$ resonance. To 0th order the NEXAFS resembles the unoccupied density of states (neglecting the light polarization, excitonic, and
FIG. 2. (Color online) (a) ARPES intensities around the K point of the Brillouin zone of graphene intercalated with one monolayer Au for increasing potassium doping. The last viewgraph (bottom right) denotes the photoemission intensities of graphene intercalated with potassium in between the graphene/Au interface. (b) Energy dispersion curves taken at the K point for the six doping steps in (a) with the shift of $E_F$ as indicated. For the doped graphene, the two peaks correspond to the $\pi$ and $\pi^*$ bands. Upon intercalation of potassium between graphene and Au, the value of the gap between $\pi$ and $\pi^*$ bands doubles.

matrix element effects). A comparison with the calculated DOS is therefore key to assign the measured feature at the $\pi^*$ resonance to an electronic state. To that end we performed tight-binding (TB) calculations of the DOS of disordered H-graphene using the equation-of-motion method for various values of H coverages. This method allows us to treat large systems with adjustable hydrogen coverages. In this case we chose graphene with 180 000 C atoms for the calculations. The TB parameters we used were $\gamma = -2.7$ eV for the C-C interaction energy, $\gamma_i = -5.7$ eV for the C-H hopping energy, and $\epsilon_i = -0.2$ eV for the on-site energy of the hydrogen orbital. In Fig. 1(d) we depict the calculated density of states of C$_n$H for various C/H ratios in agreement with previous calculations of disordered graphene. It can be seen that with increasing H/C ratio, a new electronic state appears in between the $\pi$ and $\pi^*$ bands at the Fermi energy. Looking at Fig. 1(c) the additional shoulder at the $\pi^*$ resonance also appears close to the onset (the edge of the conduction band in Fig. 1 is located at 284.2 eV). It is therefore likely that the additional shoulder in NEXAFS is a fingerprint of a hydrogen-derived midgap state. Such midgap states have been theoretically predicted but have not been observed so far in graphene. Their existence may have implications for superconductivity comparable to the case of boron-doped diamond as well as magnetism and optical properties of H-graphene.

Therefore, we turn to an alternative strategy which comprises the doping of graphene with additional electrons. This results in a shift of the Fermi level with respect to the Dirac point so that the hydrogen midgap state becomes populated with electrons and should be observable with ARPES. A well-known approach from graphite is the intercalation of potassium, since alkali metals donate their electrons without forming covalent bonds which might influence the hydrogenation procedure. Figure 2(a) shows ARPES spectra of monolayer graphene with an increasing potassium coverage. The first viewgraph (upper left) in Fig. 2(a) corresponds to undoped graphene and one can see the $\pi$ band touching $E_F$ at the K point in the Brillouin zone. Clearly, the Dirac point shifts away from $E_F$ to higher energies with increasing potassium coverage up to a final value of $\sim 1$ eV. It has been put forward that the small gap in the spectral function at the K point originates from the Au superstructure which breaks the AB symmetry. Warming up the sample to RT and cooling to 25 K again yields a completely different picture shown in the last panel (bottom right) of Fig. 2(a): A much larger separation between $\pi$ and $\pi^*$ at the K point is observed. We attribute this to the intercalation of potassium atoms into the graphene/Au interface, consistent with the behavior of other metals (Au, Ag, Fe, Cu...) that readily intercalate in between the interface of graphene and the Ni substrate. The intercalation at RT is also consistent with the reported 100 K temperature limit above which potassium ions on a graphene sheet become mobile. Upon potassium intercalation the gap between $\pi$ and $\pi^*$ bands increases which could be explained by the fact that the Au lattice forces the potassium ions to positions which distort the graphene lattice and further break the AB symmetry of the carbon atoms. The energy dispersion
curves (EDCs) at the $K$ point are shown in Fig. 2(b) for the potassium-doping steps. The $n$ doping does not significantly change the separation between the $\pi$ and $\pi^*$ bands. However, upon potassium intercalation the energy separation increases from 400 meV to 800 meV. We purposefully induce this “gap” for a direct observation of the hydrogen acceptor level which we will discuss below.

In Fig. 3(a) we present ARPES spectra of a hydrogenation series that was performed on fully $n$-doped graphene. In the potassium intercalated graphene with the larger gap between $\pi$ and $\pi^*$, the graphene layer provides a buffer between potassium ions and H atoms on top of graphene which efficiently prevents chemical bonding. From the hydrogenation series in Fig. 3(a) we observe (i) a general broadening of the spectra, (ii) a shrinkage of the Fermi surface pointing towards hole doping, (iii) that the gap between $\pi$ and $\pi^*$ increases with hydrogenation, and most importantly, (iv) that a new state appears within this gap. The new state is almost dispersionless and its ARPES intensity increases as the new electronic midgap state at 1 eV that we have found using ARPES. For a complete understanding of the origin of this new state, we project the DOS on the $C 2p_z$ and $H 1s$ orbitals. We consider $C 2p_z$ from the hydrogenated site up to the third-nearest neighbor. Figure 3(d) depicts the projected DOS for the midgap state. Interestingly, the contribution of the $2p_z$ orbital of the same sublattice of the hydrogenated C atom is almost zero. From the projected DOS it is evident that the midgap state is made up largely from $H 1s$ orbitals and the $C 2p_z$ orbitals from the neighboring lattice site. In Fig. 3(d) these two atoms are indicated by $H$ and $C_1$. The supplementary figures S4 depicts a realistic DFT calculation of hydrogenated graphene on a K/Au substrate and indicates that the midgap state is robust against substrate interactions.

In Fig. 3(b) we depict an analysis of the spectral functions from Fig. 3(a) of hydrogenated $n$-doped graphene at the $K$ point which fortifies our findings. Already at ratios of $H/C = 0.9\%$ the energetic distance between the $\pi$ and $\pi^*$ bands increases from 0.8 eV to 1.6 eV which we attribute to the partial $sp^3$ hybridization as discussed previously. The EDCs integrated over all $k$ values for $H/C = 2.1\%$ from last panel of Fig. 3(a) unambiguously visualizes the new electronic state between 0.7 eV and 1.6 eV [top curve of Fig. 3(b)]. The lack of a dispersion of this feature can be clearly seen when comparing the integrated EDC with the single EDC taken at the $K$ point. Whereas the shape of the midgap state is nearly identical in both curves, only the intensity of this state is increased when integrating the EDCs in $k$ space. Figure 3(c) shows DFT calculations of the band structure of H-graphene HC$_{32}$ in agreement with previous calculations, which is comparable to that measured for the highest $H/C$ ratio of 2.1%. To account for the $n$ doping from potassium atoms, we applied a rigid shift by 1 eV. Clearly, there is an almost dispersionless shallow acceptor level also present in the calculation which we identify as the new electronic midgap state at $\sim -1$ eV that we have found using ARPES. For a complete understanding of the origin of this new state, we project the DOS on the $C 2p_z$ and $H 1s$ orbitals. We consider $C 2p_z$ from the hydrogenated site up to the third-nearest neighbor. Figure 3(d) depicts the projected DOS for the midgap state.
Our present data suggest that the midgap state always exists in hydrogenated graphene, but its position with respect to \( E_F \) determines whether it is observable by ARPES which probes only occupied states or absorption spectroscopies such as NEXAFS. Only if \( E_F^* \) is above the energetic position of the midgap state (denoted by \( \sigma_0^* \)) can it be directly observed in ARPES. In this case, the midgap state likely accepts electrons from the \( \pi^+ \) band of graphene denoted by arrows (see the right panel).

Finally, we investigate impurity band formation and electron localization in H-graphene which governs its transport and optical properties. On the one hand, a minimum concentration (Mott criterion) is needed for band formation, but on the other hand, too much disorder can induce electron localization in both the \( \pi \) electron bands and the midgap state. A very low H impurity concentration leads to midgap energy levels corresponding to bound states. The radius of such bound states is given by \( a_B = e^2/(2\varepsilon E_B) \) where \( \varepsilon \) is the relative dielectric constant of the host material (in this case graphene) and \( E_B \) is the binding energy of the acceptor level. The Mott criterion\(^1\) for the formation of a metallic band from such discrete levels states that the concentration \( n_\varepsilon \) of the impurities must be high enough to satisfy \( n_\varepsilon \approx 1/4 \).

Assuming the H/C ratio to be \( \eta \), the critical concentration will be given by \( n_\varepsilon = 0.007(\varepsilon E_B)^2 \). In our case the position of the impurity band with respect to \( E_F \) is given by \( E_B \approx -1.0 \) eV, and with \( \varepsilon \approx 2 \) the critical concentration needed for metallization is \( \eta_\varepsilon \approx 1.4\% \). Therefore when the concentration of H atoms is \( \approx 1\% \), we expect the formation of an impurity band. However, since the position of H adsorbates is random (see supplementary information), the question of the localization of impurity band states arises. To answer this question, we use the KPM method to calculate the typical electron density of states, wherein the vanishing of this DOS indicates localization.\(^16\) In Fig. 5 we have depicted the average DOS and the typical DOS for H-graphene within a TB model.\(^11\) Such a TB model, although ignoring the effect of the substrate, is consistent with our ARPES data for the nonhydrogenated samples which show perfect Dirac cone dispersion. When the concentration of H atoms is very low, the model results in an effective Hamiltonian for C atoms only, with an energy-dependent on-site potential.\(^11\) However, when the H impurity concentration is comparable to \( n_\varepsilon \), in addition to the energy-dependent random potential (Anderson type), the hydrogenic wave functions will have substantial overlap to give rise to an impurity band. Moreover, for moderate values of the on-site disorder \( W \approx \gamma \), where \( W \) is the range of the on-site energies and \( \gamma \) is the C-C hopping energy, a mobility edge emerges in the conduction and valence bands.\(^16\)

The acceptor band survives the randomness regarding the position of the H atoms, and the states in the center of the acceptor band remain extended and a localization of these states is not expected. As can be seen in Fig. 5, a vanishing of the typical DOS indicates that the states close to the edges of the impurity band get localized, but due to the substantial bandwidth, states in the middle of the band remain extended. Therefore, although the impurity band has no dispersion as indicated by the ARPES measurements of the spectral function, it nevertheless has a sufficiently large bandwidth, so as to provide enough kinetic energy to enable conduction within this band. For very low concentrations, a localization of charge carriers in this impurity state would be expected, as the right and left mobility edges of the impurity band are expected to merge when the impurity band becomes narrow enough.

### IV. CONCLUSIONS

In conclusion, we have found a new electronic state in H-graphene that is located between the \( \pi \) and \( \pi^+ \) bands. For undoped H-graphene this state is energetically situated within the gap around \( E_F \) and is accessible with absorption spectroscopies such as NEXAFS. In the case of \( n \)-doped H-graphene the midgap state becomes available for electrons and directly observable with ARPES since it is then situated below \( E_F \). Therefore, the H impurity band likely acts as an electron acceptor level which provides the possibility to control the electron concentration in H-graphene via the H/C ratio. An estimation of the Mott criterion and a calculation of the typical DOS suggests that above \( H/C \approx 1\% \) and below \( H/C \approx 6\% \), the acceptor level can form an extended impurity band. DFT...
calculations of the DOS show this new band to be largely composed from H 1s orbitals. Hence the new electronic state we found is expected to give rise to metallic conduction when the chemical potential is tuned to cross the impurity band. Furthermore the questions of a spin splitting of the impurity band and the magnetic properties of H-graphene arise as has been suggested by theoretical calculations. Further studies including spin-resolved methods might be necessary to address these open questions and gain more detailed insights into the nature of this new electronic state in H-graphene.

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