Dynamical effects in x-ray absorption spectra of graphene and monolayered h-BN on Ni(111)

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We present first-principles calculations of x-ray absorption spectra of graphene and hexagonal BN monolayer on the Ni(111) substrate. Including dynamical core-hole screening effects according to the theory of Mahan–Nozières–de Dominics (MND) results in an overall good agreement with previously published experimental data and our new observations. This approach provides a unified first-principles description of the electronic structure and direct excitations in the sp2-bonded materials on metal surfaces and a better insight into the dynamics of screening effects. We demonstrate in particular that the observed spectral features of graphene and hexagonal BN can be well reproduced with the MND theory, and that they are determined by a delicate balance between initial and final-state effects.

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X-ray absorption spectroscopy (XAS) plays an important role in revealing the electronic structure of the sp2-bonded layered materials such as graphite and hexagonal boron nitride (h-BN). The element and symmetry selectivity of XAS makes it a convenient tool for probing the B 2p, C 2p, and N 2p density of states (DOS) in h-BN (Refs. 1–7) and graphite.8–12 However, the influence of the core hole on the spectral shape hampers a straightforward interpretation of the spectra, as was recognized already in the early works.2,3,9,13 The near-edge absorption fine structure (NEXAFS) at the B 1s, C 1s, and N 1s thresholds in h-BN and graphite is affected differently by the corresponding core holes, due to the different core-hole screening. Therefore, a correct theoretical description of all XA spectra in h-BN and graphite is difficult from an initial or final-state approach, as well as from a transition state model, because the degree of screening is very different and unknown a priori. Moreover, it can be necessary to assume a co-existence of good and poor screening and to use different theory levels even for one and the same spectrum, e.g., C 1s NEXAFS in bulk graphite.14

The problem of a unified theoretical description of the NEXAFS in graphite and h-BN is especially severe for the π* excitations due to the strong differences in their life time. As shown in Fig. 1, the full width at half maximum of the π* resonance (peak A) in bulk materials (black curves) varies from 0.3 eV for the B 1s−1 π* excitation to 1.1 eV for the C 1s−1 π* excitation and further to several eV for the N 1s−1 π* excitation. Since the symmetry of the potential for the core-excited electron is the same in all three cases, the strong variation in life time of the π* excitation is exclusively due to the variation in the electron density on the absorbing site resulting, in particular, in different screening behavior. The differences in the core-hole lifetime account for the different decay dynamics. The relative rate of the nonradiative participator Auger decay is about 30% for the B 1s−1 π* excitation in h-BN,7 about 2% for the C 1s−1 π* excitation in graphite,11 and below 0.5% for the N 1s−1 π* excitation in h-BN,7 as determined by resonant photoemission.

Thus, large variations in the screening characteristics between the B 1s, C 1s, and N 1s pre-edge excitations in the sp2-bonded materials has hitherto hindered the formulation of a unified theoretical treatment equally suitable for all three situations. In other words, the initial-state, final-state, and dynamical effects contribute to the spectral shape in different and unknown proportions. A well-defined and controlled modification of the electronic structure of graphite and h-BN can be easily performed by forming a single atomic layer of these materials on a reactive transition metal surface. The chemisorption of monolayer graphite (MG or graphene) and h-BN on the lattice-matched and chemically active Ni(111) substrate is among the most popular interfaces studied in the

FIG. 1. (Color online) Experimental B 1s, C 1s, and N 1s NEXAFS spectra from monolayer graphite and h-BN adsorbed on Ni(111) compared with the corresponding spectra of bulk HOPG and h-BN. The energy scales are aligned at the position of peak A in the spectra of bulk materials. The angle between the surface normal and the polarization vector of incident radiation is 50° for all spectra.
past. The electronic structure of both graphene and h-BN on Ni(111) is perturbed in a similar manner due to the Ni 3d—MG (h-BN) π orbital mixing.15–18

In this paper we systematically treat the process of x-ray absorption in MG/Ni(111) and h-BN/Ni(111) at the B 1s, C 1s, and N 1s edges by initial-state, final-state, and dynamical calculations, and compare the results with the B 1s, C 1s, and N 1s NEXAFS spectra. The experimental data for h-BN/Ni(111) were measured previously,17 while for the MG/Ni(111) system only the electron energy loss spectra (EELS) were reported so far.19 First-principles calculations were performed in the past to explain both EELS in MG/Ni(111),19 and NEXAFS spectra in h-BN/Ni(111).20 Although the agreement with experiment was found to be rather good, only the static effect of the core hole on the spectra could be revealed in these calculations. Here we aim at providing a unified description of the x-ray absorption process in all three cases, and demonstrate that the overall spectral evolution of XAS in graphite and h-BN is going from bulk materials to monolayers.

The formation of exactly one graphite ML was checked by the single-component shape of the C 1s photoelectron peak A, which is widely used as a visible effect is a reduction in the energy separation between features is needed, and is the focus of this study. Another formalism described in Refs. 14 and 28. The energy range of the Green’s functions and MND calculation was from −2 Ry down to 0.02 Ry above the Fermi level, served as normalization coefficients (i indexes the natural orbitals). In the second step we transformed all \( \langle lms | k_\nu (k_\mu l' m' s') \rangle \) matrices into the natural orbital basis, renormalized their elements by 1/\( \sqrt{n_n} \), and transformed back into the \( \langle lms \rangle \) basis. After this transformation, the energy-integrated Green’s function matrix equals to the unit matrix. These normalized Green’s function matrices serve as an input into an efficient MND implementation based on formalism described in Refs. 14 and 28. The energy range of the Green’s functions and MND calculation was from −2 Ry up to 3 Ry (zero at Fermi level) with a step of 0.005 Ry and a smearing of 0.01 Ry.

The calculated spectra are summarized in Figs. 3 and 4. In these figures we compare the experimental spectrum (black)
The local electronic structures of boron and nitrogen in the presence of the core hole. This shows that the local screening electrons available for screening of the core hole.

The situation of the nitrogen edge is more complicated. Neither the initial nor final-state calculation matches the measured spectra. Although the origin of the main experimental structure at 4 eV (the $\pi^*$ band shifted upwards by the orbital mixing with the Ni $3d$ states) can be reproduced by both initial-state and final-state calculations, the shape of the N 1s absorption onset is clearly incorrect. Besides, the $\sigma^*$ bands are largely dominating the electronic structure at 6–12 eV and they decay too quickly at higher energies. The situation is dramatically improved by the MND calculation, showing the significance of the dynamical screening pro-
cesses on the N 1s spectral shape. This improvement is facilitated by the gapless nature of the ground-state electronic structure of chemisorbed h-BN due to a considerable density of gap states of primarily N 2p(π*) character induced by the orbital mixing at the h-BN/Ni(111) interface.29 For gapless systems the MND approach is known to work especially well, because the core-hole relaxation is crucial there due to the high probability of creating shake-up pairs with low energy.

The case of graphene on Ni(111) is somewhat complicated by the co-existence of two chemically nonequivalent C atoms, see Fig. 2, with different number of nearest neighbor Ni atoms.19 This causes differences in their local electronic structure, particularly in the π DOS. We show in Fig. 4 spectral features projected on both atom types, for the spectra calculated at the initial-state, final-state, and MND theory levels. Neither initial-state nor final-state theory reproduces the experimental curve properly, whereas the MND spectrum provides a significant improvement of the spectral shape. The experimentally observed two-peak structure at the K-edge onset is very well reproduced and the MND σρ peak at 7 eV has now a proper relative magnitude (note the large overestimation of its magnitude in both initial and final states). The hybridization-induced peak at 2 eV is present mainly in the MND spectrum originating from the carbon atoms on top of the fcc-hollow sites. These atoms are somewhat closer to the substrate19 and their pz orbitals can overlap with the lobes of the Ni 3d(t2g) orbitals directed toward these sites. Therefore, despite their larger absolute distance from the nearest Ni atoms (compared to the C on top), their local electronic structure is more influenced by the substrate. Note that both initial-state and final-state calculations provide only a qualitative description of these phenomena, while the explicit inclusion of electron correlations into the calculation of transition probabilities results in the correct description of the spectral shape.

In summary, we have succeeded in reproducing all experimental XAS spectra of graphene and h-BN monolayers interacting with the Ni(111) substrate, using a parameter-free first principles method, which explicitly takes into account the dynamics of the core-hole screening. These dynamical screening effects are particularly important for the nitrogen and carbon K-edges, due to the high density of the N 2p and C 2p states in the vicinity of the Fermi level. For the boron K-edge the MND theory results are similar to those of a more simple static final-state calculation, as a consequence of the poor core-hole screening. In general, our study shows that the real dynamics of the core-hole screening is crucial for correct description of x-ray absorption processes in this class of layered materials and their contacts with substrates. We also show that the new spectral feature, A′, is due to a combination of dynamical core-hole screening and the hybridization between the π orbitals and the Ni 3d states.

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