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Exciton energy-momentum map of hexagonal boron nitride

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Understanding and controlling the way excitons propagate in solids is a key for tailoring materials with improved optoelectronic properties. A fundamental step in this direction is the determination of the exciton energy-momentum dispersion. Here, thanks to the solution of the parameter-free Bethe-Salpeter equation (BSE), we draw and explain the exciton energy-momentum map of hexagonal boron nitride (h-BN) in the first three Brillouin zones. We show that h-BN displays strong excitonic effects not only in the optical spectra at vanishing momentum \( q \), as previously reported, but also at large \( q \). We validate our theoretical predictions by assessing the calculated exciton map by means of an inelastic x-ray scattering (IXS) experiment. Moreover, we solve the discrepancies between previous experimental data and calculations, proving then that the BSE is highly accurate through the whole momentum range. Therefore, these results put forward the combination BSE and IXS as the tool of choice for addressing the exciton dynamics in complex materials.

I. INTRODUCTION

The response of materials to electromagnetic fields is determined by electronic excitations that are strongly influenced by electron-hole (e-h) interactions. In particular, the e-h attraction leads to the formation of excitons, which are a fundamental aspect in the functionality of many optoelectronic devices, as excitons can propagate in materials carrying excitation energy that can be transformed and exploited by different means. Excitons can be identified in electronic spectra as sharp peaks within the band gap of insulators, or, beyond the band gap, as spectral intensity enhancement towards lower energies with respect to a non-interacting theoretical picture. A reliable description and analysis of those two-particle correlation effects is therefore the key to understand materials’ dielectric properties, guide the realisation of new experiments and foster the development of new technological applications.

Nowadays, the ab initio solution of Bethe-Salpeter equation (BSE) represents the state-of-the-art method to obtain spectra in very good agreement with experiments in a large variety of materials. These theoretical achievements have mainly focused on optical absorption and electron energy-loss spectroscopy (EELS) spectra for vanishing momentum transfer \( q \rightarrow 0 \). However, judging whether a theoretical approach captures fully the physics of the electron dynamics requires stringent tests that are offered by the measurement of the full dynamic range of the relevant variables (momentum and energy). This assessment is now possible thanks to the spectacular progress of inelastic x-ray scattering (IXS) experiments, both in the resonant (RIXS) and non-resonant (NRIXS) conditions. They allow one to probe electronic excitations at finite momenta \((q)\) with a resolving power that has improved by orders of magnitude in the last two decades. The challenge for theory is hence the first-principles description of the full electron dynamic, which is well beyond the sole simulation of optical absorption spectra. The investigation of the energy-momentum dispersion of elementary excitations (excitons, plasmons, etc.) provides fundamental information on the way they propagate in materials. Moreover, spectroscopic features measured at larger \( q \) allow for the study of excitations that in real space occur on shorter interatomic scales and/or are not visible in optics because they are dipole forbidden (see e.g. Refs. [1](10,11,12]).

Recently, some of us have shown that these ambitious goals are within reach, thanks to the extension of the BSE to describe e-h excitations carrying a finite momentum \( q \). In the case of a prototypical wide-gap insulator such as lithium fluoride, the dynamic structure factor \( S(q, \omega) \) was obtained in excellent agreement with accurate NRIXS data. However, in contrast to LiF, in hexagonal boron nitride (h-BN), a layered material that is the insulating counterpart of graphite, a recent comparison between BSE calculations and NRIXS data revealed a mismatch at high momentum transfers. The experiment displays a sharp peak at \( \sim 7 \) eV whose origin has still to be understood, as pointed out in Ref. [14]. Moreover, despite the accurate BSE results in LiF, a recent study based on a simplified exciton kinetic kernel model raised doubts about the capability of the ab initio BSE in general to address the issue of the exciton band structure. Therefore new questions arise: Is LiF only a fortunate case for BSE? For spectra at finite \( q \) should we hence go beyond the standard BSE implementation that has been successfully applied for optics? Or, on the contrary, is the sharp peak measured in h-BN an artifact of the NRIXS experiment? Finally, is the BSE...
the appropriate method in general to study the exciton band structure of complex materials.

In the present work, we demonstrate that it is possible to reconcile theory and experiment if the microscopic details of the screened Coulomb e-h interaction are explicitly taken into account in the calculations. We show that in h-BN excitonic effects not only produce a redshift of the peaks, as observed in Ref. [19] but also lead to important spectral shape redistributions. We reproduce all the details of the NRIXS spectra measured in Ref. [19] including the 7 eV peak at large $q$, and we consistently explain the appearance of excitonic features at various $q$ as a multiple manifestation of the same large joint density of states (JDOS). Moreover, by calculating the entire map of $S(q,\omega)$ in the first 3 Brillouin zones of h-BN, we identify new “hot spots” in dynamics, where the probability for creating an exciton is the largest. These predictions are fully confirmed by the new NRIXS experiment that we have performed at the beamline ID20 of the European Synchrotron Radiation Facility (ESRF) in Grenoble (France). Therefore these new results demonstrate that the combination of first-principles BSE calculations and accurate NRIXS experiments is a very powerful tool to explore and understand the exciton dynamics.

II. THEORY

The spectrum obtained in an NRIXS experiment is proportional to the dynamic structure factor:

$$S(q,\omega) = -\frac{q^2}{4\pi^2 n} \text{Im} \epsilon_M^{-1}(q,\omega),$$

where $n$ is the average electron density. The inverse of the macroscopic dielectric function $\epsilon_M^{-1}$ can be expressed as:

$$\epsilon_M^{-1}(q,\omega) = 1 + \frac{8\pi}{q^2} \sum_{\lambda} \left| \sum_{t} A^t_{\lambda}(q) \tilde{\rho}^{t}(q) \right|^2 \frac{\omega - E_{\lambda}(q) + i\eta}{\omega - E_{\lambda}(q)},$$

where in the Tamm-Dancoff approximation (TDA) the sum is over valence-conduction ($v-c$) transitions $t$, the oscillator strengths are $\tilde{\rho}^{t}(q) = \langle \phi_{v,k-q} | e^{-iqr} | \phi_{c,k} \rangle$, $k$ and $q$ are in the first Brillouin zone, and $q = q + G$ is the measured momentum transfer with a reciprocal-lattice vector $G$. The BSE can be cast into an effective two-particle Schrödinger equation. The exciton hamiltonian $H_{\text{exc}}$: $\langle t | H_{\text{exc}} | t' \rangle = E_{\delta}(t,t') + \langle t'| v_e - W | t' \rangle$ contains the quasiparticle e-h transition energies $E_{\delta}$ calculated in the GW approximation, and the matrix elements in the transition basis of the bare Coulomb interaction $v_e$ and the statically screened Coulomb interaction $W = \epsilon^{-1} v_e$, which describes the e-h attraction (i.e. gives rise to excitons) and is here obtained in the random phase approximation (RPA). We have calculated the ground state of h-BN within the local density approximation (LDA) of density functional theory (DFT) using norm-conserving Troulliers-Martin pseudopotentials in a plane wave approach with an energy cutoff of 30 Hartree. In order to approximate GW quasiparticle energies, following we corrected the LDA band structure with a scissor operator of 1.98 eV for the band gap and stretching the valence bands by 5%. The BSE spectra at finite $q$ are obtained from the EXC code using 25 bands and a $18 \times 18 \times 4$ $k$-point grid for calculations with $q$ parallel to the hexagonal plane and with a $12 \times 12 \times 8$ $k$-point grid for calculations with $q$ perpendicular to it.

III. EXPERIMENT

In the NRIXS experiment the incident x-ray beam from three undulators was monochromated to an energy of 7.5 keV by a combination of a Si(111) double crystal and a Si(311) channel-cut. The beam was focused to a spot of $\approx 10 \ \mu m \times 20 \ \mu m$ ($V \times H$). The spectrometer used a diced Si(533) analyzer crystal in the Johann geometry with a Rowland circle diameter of 2 m. The active diameter of the analyzer crystal was 80 mm, yielding a momentum-transfer resolution of $\sim 0.15 \ A^{-1}$. The energy resolution was 200 meV (FWHM). The scattering plane was vertical, i.e., perpendicular to the plane of linear polarization of the incident and scattered radiation. The detector was based on a Timepix chip, with a pixel size of 55 $\mu m$ and enabling us to use the dispersion compensation algorithm. The analyzer Bragg angle was fixed at $87^\circ$ and the energy-loss spectra were measured by scanning the incident-photon energy. The h-BN single crystal was a colorless and transparent platelet with a thickness of 0.5 mm and lateral size of 2.1 mm, which was produced by a high-pressure and high-temperature (HPHT) method using a barium-related solvent system as reported in. The samples were aligned using x-ray diffraction on the beamline.

IV. RESULTS

In layered h-BN the electronic states can be classified according to the even ($\sigma$) and odd ($\pi$) parity with respect to the single BN sheet. Optical and EELS spectra in the low-energy range, where excitonic effects are more relevant, are determined by $\pi - \pi^*$ transitions that are mainly visible for in-plane $q \rightarrow 0$. The weak screening of the Coulomb e-h interaction, in concomitance with a JDOS peak due to vertical transitions between parallel bands (see Fig. 2 in the HKML plane of the Brillouin zone), in the $q \rightarrow 0$ spectra gives rise to prominent exciton peaks with large binding energies.

Fig. 1 shows the dynamic structure factor $S(q,\omega)$ in the energy range of the $\pi - \pi^*$ transitions at three finite momentum transfers: $q = A$, K, and M, which are located at the boundaries of the first Brillouin zone (see
Fig. 1. Dynamic structure factor at three Brillouin zone boundaries $q = A, K,$ and $M$ calculated in the GW-RPA and from the solution of the BSE compared to the NRIXS data from Ref.\textsuperscript{19}.

As noticed in\textsuperscript{19} the e-h attraction induces a redshift of the peaks, which is larger for the in-plane $q$ directions at $K$ and $M$ (redshift of 1.5 eV) than for $q$ at $A$ (redshift of 0.4 eV), i.e. along the direction perpendicular to the BN layers, implying an anisotropic effect of the e-h interaction.\textsuperscript{19} This is evident from the comparison of the BSE spectra with the GW-RPA results, obtained starting from the GW band structure and neglecting the e-h attraction $W$ in the exciton hamiltonian $H_{exc}$. At variance with\textsuperscript{19} where the shift was inferred from the adjustment of the calculations to the experimental spectra, the anisotropic redshift of the spectra is here the direct outcome of the BSE calculations that result in very good agreement with NRIXS. Our simulations of the experimental spectra, being free from any adjustable parameters, allow us to additionally observe that excitonic effects also induce an important redistribution of the spectral weight towards lower energies with respect to the non-interacting e-h picture.

At $q = M$ the two main peaks are located at $\sim 8$ and $\sim 12$ eV. At the $K$ point, the first peak at $\sim 6-7$ eV appears as a shoulder of the second one. In both cases the 12 eV peak originates from non-vertical (i.e. $k \rightarrow k' = k + q$) $\pi - \pi^*$ transitions that disperse isotropically as a function of in-plane $q$, shifting the energy of the peak from 9 eV at $q \rightarrow 0$ to 12 eV in both $\Gamma K$ and $\Gamma M$ directions. The first peak, instead, is due to a peculiar property of the hexagonal Brillouin zone, see Fig. 4(a). For $q = M$ it derives from vertical e-h transitions between $k$ points belonging to the ML line in the band structure (see Fig. 2) (analogous, for $q = K$, is the HK line). These are the same vertical $\pi - \pi^*$ transitions with large JDOS that are at the origin of the tightly bound exciton in the spectra at $q = 2.5 \text{Å}^{-1}$ is the $M'$ point (see text).

Having clarified the diverse role of excitonic effects at various $q$, we are ready to discuss what happens along the $\Gamma K$ direction, where for $q > 2.0 \text{Å}^{-1}$ a peak appears in the NRIXS spectra at $\sim 7$ eV that was unexplained in Ref.\textsuperscript{19}. The spectra that we have calculated for different momentum transfers along $\Gamma K$ demonstrate that the BSE is actually able to reproduce the 7 eV peak that is measured in the NRIXS experiment\textsuperscript{19} (see Fig. 3). Moreover they show that the peak has indeed an excitonic character as it is completely absent in the GW-RPA calculations. In order to better understand the origin of this excitonic peak one has to note that it has the largest intensity at $q = 2.5 \text{Å}^{-1}$. This momentum transfer along the $\Gamma K$ direction is in fact another M point, which is located at the boundary of the third Brillouin zone and

![Fig. 2](image2.png)

**Fig. 2.** (a) GW band structure of hBN in the (b) first Brillouin Zone with the irreducible part delimited by yellow lines and shaded.
FIG. 4. (a) Color map of $S(q, \omega)$ calculated from the BSE for fixed $\omega = 7$ eV and for all $q$ spanning the first 3 Brillouin zones with $q_z = 0$. To enhance its visibility, $S(q, \omega)$ in the first BZ has been magnified by a factor of 3. The high-symmetry points relevant for the discussion in the main text are explicitly indicated (in parenthesis those belonging to the $q_z = 0.5$ plane). Adding $q = \Gamma M$ (solid violet line) to $k = M (L)$ one obtains another $k' = k + q = M (L)$ (see dashed violet line). Equivalently adding $q = \Gamma K$ (solid red line) to $k = K (H)$ one obtains again $k' = K (H)$ (see red dashed line). (b) Comparison between calculated and measured $S(q, \omega)$ for $q$ along the $\Gamma M$ direction.

which we call $M'$. We can also immediately recognise why excitonic effects are again particularly strong at this momentum transfer $M'$. The exciton eigenvalues $E_{\lambda}(q)$ and eigenvectors $A_\lambda(q)$ at $M'$ must be the same as at the $M$ point in the first Brillouin zone since the two points just differ by a reciprocal-lattice vector $G$. However the oscillator strengths $\rho^0(q)$ are generally different at different $q$ and this explains why the spectra at $M$ and $M'$ do not entirely overlap. In particular, at $q = M'$ there is a prominent peak at 7 eV emerging from a featureless plateau at higher energy, while at $q = M$ a double-peak structure was observed in the same energy range.

We have therefore found that the solution of the BSE, when the screening of the Coulomb interaction is explicitly calculated at the RPA level rather than obtained from model dielectric functions, is able to reproduce and explain the NRIXS data. This is confirmed by the direct comparison between the calculated and experimental spectra (see Appendix). The agreement is excellent in all considered cases. Moreover, having understood that strong excitonic effects for the peak at 7 eV should be expected whenever $\pi - \pi^*$ transitions give rise to an intense JDOS, we can forecast other “hot spots” with large intensity in the dynamic structure factor, beyond what has been experimentally detected in Ref. 19. In Fig. 4(a) we thus plot a color map of $S(q, \omega)$ for a fixed $\omega = 7$ eV and for all $q$ spanning the first 3 $q_z = 0$ Brillouin zones. In this manner we can easily identify two other remarkable points along the $\Gamma M$ direction, located between $q = M$ and the vertex of the hexagon $q = \Gamma'$, which differs from the $\Gamma$ point by a reciprocal-lattice vector $G$.

In order to confirm these predictions and assess the whole theoretical map of $S(q, \omega)$ that has been obtained from the BSE calculations, we compare the theoretical results to those obtained from our NRIXS experiment in Fig. 4(b). While the GW-RPA results are totally different from the experiment, the excellent agreement between the BSE results and the NRIXS data proves the predictive power of the BSE. In the spectra the largest intensity is found at $q = M'$, displaying two prominent peaks deriving again from the large JDOS of vertical $\pi - \pi^*$ transitions. The first peak, visible only close to $q = \Gamma'$, matches the main exciton peak at the onset of the spectrum at $q \to 0$. The second peak at $q = \Gamma'$, which is not noticeable at $q \to 0$, instead evolves continuously from the first peak located at $\sim 8$ eV at $q = M$. These results show that inspecting spectra at large $q$, thanks to the variation of the oscillator strengths with $q$, can reveal formation of excitons that are hidden in corresponding optical spectra. Moreover, they evidence how the combination of NRIXS and BSE provides the means for their detection in the whole energy-momentum range, allowing one to obtain the full exciton band structure.

V. CONCLUSIONS

In summary, by solving the discrepancies between previous experiments and calculations, we have successfully established the energy-momentum map covering the first 3 Brillouin zones of the dynamic structure factor $S(q, \omega)$ of h-BN, a prototypical layered insulator. We have shown that excitonic effects in h-BN are strong also at large momentum $q$. They are essential to interpret and understand the spectra (including previous unexplained features) and cannot be neglected remaining at a level of theory corresponding to the RPA. We have explicitly proved that such a theoretical map based on BSE is crucial in order to guide the experimental exploration of the electronic dynamics, as new IXS measurements
FIG. 5. Dynamic structure factor along three high symmetry lines in the first Brillouin zone ΓA, ΓK, and ΓM calculated in the GW-RPA (dashed lines) and from the solution of the BSE (solid lines), compared to the NRIXS data (dotted lines) from Ref.10. The distance from Γ to the borders of the 1st BZ along each direction is indicated in the labels.

have fully validated the theoretical predictions. Therefore these case-study results promote the ab initio solution of the BSE, which has been already successfully applied in optics, as an accurate and predictive method also to investigate the charge dynamics and the exciton band structure for a wide range of materials and technological applications.

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Appendix: Dynamic Structure Factor in the First BZ

We report in Fig. 5 the comparison between the calculated and experimental spectra along three high symmetry lines ΓA, ΓK, and ΓM in the first Brillouin zone.

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