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Citation: The Journal of Chemical Physics 126, 214306 (2007); doi: 10.1063/1.2741524
View online: http://dx.doi.org/10.1063/1.2741524
View Table of Contents: http://aip.scitation.org/toc/jcp/126/21
Published by the American Institute of Physics
Photoabsorption spectra of boron nitride fullerenelike structures

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(Received 23 February 2007; accepted 25 April 2007; published online 5 June 2007)

Optical absorption spectra have been calculated for a series of boron nitride fullerenelike cage structures B$_n$N$_n$ of sizes $n = 12–36$. The method used is a real-time, real-space implementation of the time-dependent density-functional theory, involving the full time propagation of the time-dependent Kohn-Sham equations. The spectra are found to be a possible tool for distinguishing between different boron nitride fullerene species and isomers. The trends and differences in the spectra are found to be related to the general geometry of the molecules. Comparison between local-density and generalized-gradient approximations for electron exchange-correlation functionals shows that both of them produce essentially the same spectral characteristics. © 2007 American Institute of Physics. [DOI: 10.1063/1.2741524]

I. INTRODUCTION

Like carbon, boron nitride (BN) is able to form nano-sized structures, for example, fullerene-like cages, onions, and tubes. The fabrication of BN nanotubes$^1$ and nested cages$^2$ was reported in the mid-1990s, soon after the discovery of carbon-based fullerenes and related carbon nanostructures. BN fullerenes were synthesized for the first time in 1998 by Stéphan et al.$^3$ These isoelectronic analogs of carbon structures are of interest to both theorists and experimentalists in materials science because of their potential applications in future nano- and optoelectronic devices and also as lubricants. The superiority of BN based structures over the carbon ones would result from their better thermal and chemical stabilities. In this work, the objective is the optical properties of BN fullerene cages calculated from first principles. Especially, we want to study their optical absorption spectra which can be used in identifying their structures.

The time-dependent density-functional theory (TDDFT) in its linear-response form is already a standard chemist’s tool for calculating optical spectra of small atom clusters and molecules. In contrast to the (time-independent) DFT, the TDDFT can properly treat the excitations of electronic systems (for a review see, for instance, Refs. 4–6). The implementations of the TDDFT are divided into linear-response methods and methods propagating the time-dependent Kohn-Sham equations in real time. Due to the unfavorable scaling of the linear-response methods as the electron number $N$ increases, they become ineffective for structures that consist of several dozens of atoms. Instead, the propagation of the Kohn-Sham equations becomes advantageous despite its large prefactor since it scales linearly with $N$. This method has been shown to reproduce the main measured low-energy excitations correctly with an accuracy of few tenths of eV for several kinds of systems as long as the time step is small enough and the total propagation time is adequate. It has been used, for example, for calculating the optical absorption spectra of different B$_{20}$ (Ref. 7) and C$_{20}$ (Ref. 8) isomers.

According to our knowledge, the results presented in this paper are the first TDDFT calculations for BN nanostructures. We are extending the knowledge on the yet almost unexplored optical properties of BN by means of a sophisticated and efficient method. An important aspect of our work is that the structural and spectral properties are calculated on the same footing for a representative set of different BN fullerene structures. This enables the highlighting of differences and trends.

II. STRUCTURES

The basic structural properties of the molecules studied in this work are given in Table I. The molecules are chosen such that they would form a series of energetically favorable structures of BN fullerenes. The following findings from the literature on both experimental results and structure calculations with chemical accuracy have been considered as guidelines for the present work.

It has been found computationally that fullerenelike cage structures B$_n$N$_n$ are energetically more favorable than ring structures for $n > 10$. Unlike carbon fullerenes, BN fullerenes consist purely of rings with even number of atoms. This is caused by the high-energy cost of placing two B or two N atoms next to each other. In computational studies for the fullerene structures most attention has been paid to those with high symmetry. The “magic” octahedronlike structures of B$_{12}$N$_{12}$, B$_{16}$N$_{16}$, B$_{28}$N$_{28}$, and B$_{36}$N$_{36}$ are found to be especially stable.$^{10–13}$ They consist purely of four- and six-membered rings. In contrast, for B$_{24}$N$_{24}$, the isomers with the lower $S_6$ and $S_4$ symmetries are computationally found more stable than the most symmetric isomer with the $O$ symmetry.$^{14,15}$ This is due to the high number of energetically less favorable octahedral rings in the $O$-symmetric structure. However, the differences in total energy are small, of the order of 0.1 eV/atom or less. The same kind of problem is observed in calculations for B$_{32}$N$_{32}$, the most stable struc-
The initial structures of the studied BN fullerenes were constructed based on the data in Table I and the literature presented in Sec. II. Then a geometry optimization using the SIESTA program was performed for fine tuning the geometries of the molecules.

The main work was carried out by the OCTOPUS program, a TDDFT real-time real-space code. The full time evolution of the time-dependent Kohn-Sham equations was calculated in OCTOPUS to obtain the dipole strength function, that is to say, the optical absorption spectrum.

Let us briefly summarize the essentials of the TDDFT method used in this work (for further details see, for example, Ref. 23). The system is excited from its initial ground state by applying an instantaneous electric field causing the potential $V(r,t) = -k_0 \gamma \delta(t)$, where $\gamma = x, y, z$ denotes the polarization direction and $k_0$ is the amplitude of the perturbation. This corresponds to multiplying the ground-state wave functions with an exponential $e^{ik_0 \gamma}$. Then the system is allowed to propagate over a finite period of time. The dynamical polarizability is

$$\alpha_n(\omega) = -\frac{1}{k_0} \int d^3 r \gamma \delta n(r, \omega),$$

where $\delta n(r, \omega)$ is the time Fourier transform of the deviation of electron density from the ground-state density of the system. The dipole strength function is obtained by averaging over the three spatial coordinates,

$$S(\omega) = \frac{4 \pi \omega}{c} \Im \left( \sum \alpha_n(\omega) \right).$$

Above, $c$ is the speed of light.

Calculations were carried out using two different exchange-correlation potentials: the local-density approximation (LDA) (Ref. 24) and the more advanced Perdew-Burke-Ernzerhof (PBE) generalized-gradient approximation (GGA). Calculations using even more sophisticated methods such as hybrid GGA’s, giving the correct asymptotic potential behavior, were not possible due to the limited choice of exchange-correlation functionals implemented in the

### TABLE I. Molecules studies.

<table>
<thead>
<tr>
<th>Species</th>
<th>Symm.</th>
<th>Polygons</th>
<th>Polar.</th>
</tr>
</thead>
<tbody>
<tr>
<td>$B_{12}N_{12}$</td>
<td>$T_d$</td>
<td>6$F_h$ + 8$F_k$</td>
<td>1</td>
</tr>
<tr>
<td>$B_{16}N_{16}$</td>
<td>$T_d$</td>
<td>6$F_h$ + 12$F_k$</td>
<td>1</td>
</tr>
<tr>
<td>$B_{24}N_{28}$</td>
<td>$O$</td>
<td>12$F_h$ + 8$F_k$ + 6$F_k$</td>
<td>1</td>
</tr>
<tr>
<td>$B_{24}N_{24}$</td>
<td>$S_h$</td>
<td>8$F_h$ + 16$F_k$ + 2$F_k$</td>
<td>2</td>
</tr>
<tr>
<td>$B_{28}N_{28}$</td>
<td>$S_h$</td>
<td>6$F_h$ + 20$F_k$</td>
<td>3</td>
</tr>
<tr>
<td>$B_{32}N_{28}$</td>
<td>$T$</td>
<td>6$F_h$ + 24$F_k$</td>
<td>1</td>
</tr>
<tr>
<td>$B_{36}N_{36}$</td>
<td>$T_d$</td>
<td>6$F_h$ + 32$F_k$</td>
<td>1</td>
</tr>
</tbody>
</table>

The main work was carried out by the OCTOPUS program, a TDDFT real-time real-space code. The full time evolution of the time-dependent Kohn-Sham equations was calculated in OCTOPUS to obtain the dipole strength function, that is to say, the optical absorption spectrum.

Let us briefly summarize the essentials of the TDDFT method used in this work (for further details see, for example, Ref. 23). The system is excited from its initial ground state by applying an instantaneous electric field causing the potential $V(r,t) = -k_0 \gamma \delta(t)$, where $\gamma = x, y, z$ denotes the polarization direction and $k_0$ is the amplitude of the perturbation. This corresponds to multiplying the ground-state wave functions with an exponential $e^{ik_0 \gamma}$. Then the system is allowed to propagate over a finite period of time. The dynamical polarizability is

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Calculations were carried out using two different exchange-correlation potentials: the local-density approximation (LDA) (Ref. 24) and the more advanced Perdew-Burke-Ernzerhof (PBE) generalized-gradient approximation (GGA). Calculations using even more sophisticated methods such as hybrid GGA’s, giving the correct asymptotic potential behavior, were not possible due to the limited choice of exchange-correlation functionals implemented in the...
current stable version of the OCTOPUS program. The norm-conserving Troullier-Martins pseudopotentials\textsuperscript{26} were used throughout the calculations. The double \( \zeta \) with polarization basis set was used in the SIESTA calculations. The numerical parameters in the OCTOPUS calculations were the following: time step of 0.0025\( \hbar /eV \), total propagation time of 37.5\( \hbar /eV \), spacing of real-space mesh points of 0.3 Å, and radius of wave function domain around each nucleus of 6–8 Å. The calculations correspond to zero temperature.

**IV. RESULTS**

The structure optimization resulted in bond lengths varying over the range of 1.41–1.49 Å (1.43–1.50 Å) when employing the LDA (PBE) scheme. The structures, their bond lengths, and total energies were in good accordance with the literature cited in Sec. II. Further details of the structure optimization results are not given here, as the calculation of optical spectra is not sensitive to small variations in the structure.

The spectra of all studied molecules can be divided into three energy regions. First, no excitations occur at energies below the highest occupied molecular orbital–lowest unoccupied molecular orbital gap. In the interesting region of about 5–10 eV several peaks appear, the details depending on the structure. Above that a broad feature extends up to several tens of eV’s. Most of the strength of the spectra is found here, but this high-energy region has little interest since it is more cumbersome to access experimentally. In

**FIG. 2.** (Color online) Photoabsorption spectra of three different isomers of B\(_{24}\)N\(_{24}\): \( O \), \( S_8 \), and \( S_4 \) isomers. The insets on the right-hand side show the optimized structures of the molecules.

**FIG. 3.** (Color online) Photoabsorption spectra of B\(_{30}\)N\(_{30}\) and B\(_{36}\)N\(_{36}\). The insets on the right-hand side show the optimized structures of the molecules. LDA approximation has been used.
addition, the method we used is insufficient to describe the behavior of the electrons in the high-energy region because it is dominated by ionization processes.

The results were both qualitatively and quantitatively quite similar for LDA and PBE. The number and grouping of the excitation peaks agree well and the intensities of the peaks follow similar trends. This is in accordance with other studies, for example, Ref. 27. Compared to the static DFT, where GGA clearly beats LDA in predicting structural properties, the situation is different in TD-DFT. The only systematic difference observed in this study is the slightly larger energy gaps obtained by PBE.

The spectra of the smallest studied molecules, B12N12 and B16N16, are shown in Fig. 1. B12N12 has sharp excitations at about 5.9 and 6.6 eV and a minor one around 5.2–5.5 eV. In the range of 7.5–10 eV several additional sharp peaks are observed. For B16N16 the major excitations take place around 8.5 eV, with several minor peaks in the range of 5.3–8 eV. The excitation gap is 5.0 eV (5.1 eV) for B12N12 and 5.2 eV (5.4 eV) for B16N16, according to the LDA (PBE) calculations. The excitation gaps are in good accordance with, for example, the ones calculated in Ref. 10.

For the larger molecules, the interesting 5–10 eV part of the spectra becomes fuzzier as it is observed in Figs. 2 and 3. However, some trends can be observed, for example, between the different isomers of B24N24. The symmetric O isomer exhibits sharp peaks around 7 and 9 eV, whereas the spectra of S8 and S4 isomers are flatter. This is due to the larger variety of different bond lengths and orientations in asymmetric molecules. The clearest peaks are identified around 9.5 eV for the S8 isomer and around 10 eV for the S4 isomer. The symmetric O isomer has an excitation gap of 5.0 eV, whereas the S8 and S4 isomers show clearly smaller gaps of about 4.3 eV. This difference is also explained by geometric arguments. The asymmetric isomers have a longer spatial dimension in one direction, involving longer π orbital chains. This implies a stronger delocalization of the valence electrons in that direction and thus a decrease in the gap.

According to Fig. 3, B28N28 shows a series of close peaks starting from 4.5 eV. The spectrum of B24N36 in Fig. 3 is more distinctive with larger features: the lowest peaks are observed at 5.4, 6.0–6.7, and 7.4–7.9. The excitation gaps are 4.4 eV (4.9 eV) for B28N28 and 4.5 eV (4.7 eV) for B36N36 using the LDA (PBE) scheme. Our spectrum of B36N36 based on the TD-DFT is much more complex and even qualitatively different from the one calculated on the DFT level in Ref. 16. A spectrum predicted correctly to the extent of the details is of vital importance for the identification of the cluster structures using the optical absorption spectroscopy.

V. CONCLUSION

In this work we have calculated the optical absorption spectra for several BN fullerenes in the size range of BnNn, n = 12–36. Differences between the molecules and even between different isomers are observed in the near ultraviolet energy region. This consolidates the view that the TD-DFT can be effectively used in determining and predicting the optical properties of middle-sized molecules and clusters. It has already been shown before that through the mutual interplay between calculations and experiments, the study of optical properties is a useful means of characterizing new materials. As the available computer resources expand and the TD-DFT methods advance, contributions of the TD-DFT in characterization, tailoring, or even designing new specific materials are expected to strongly increase.

ACKNOWLEDGMENTS

This research is supported by the Academy of Finland through the Centers of Excellence Program (2006–2011). CSC, the Finnish IT Center for Science, is acknowledged for providing computer resources (Project No. ttk2035).