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Published in:
Journal of Chemical Physics

Link to article, DOI:
10.1063/1.3300129

Publication date:
2010

Document Version
Publisher's PDF, also known as Version of record

Link back to DTU Orbit

Citation (APA):
Optical to ultraviolet spectra of sandwiches of benzene and transition metal atoms: Time dependent density functional theory and many-body calculations

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(Received 16 September 2009; accepted 7 January 2010; published online 28 January 2010)

The optical spectra of sandwich clusters formed by transition metal atoms (titanium, vanadium, and chromium) intercalated between parallel benzene molecules have been studied by time-dependent density functional theory (TDDFT) and many-body perturbation theory. Sandwiches with different number of layers, including infinite chains, are considered. The lowest excitation energy peaks in the spectra are characteristic of the robust bonding in these complexes. The excitation energies vary in a systematic way with the metal atoms and with the cluster size, and so these materials could be used to tune the optical properties according to specific functionality targets. The differences in the spectra could be used to identify relative abundances of isomers with different spins in experimental studies. As a salient feature, this theoretical spectroscopic analysis predicts the metallization of the infinite (TiBz)n chain, which is not the case of (CrBz)n. © 2010 American Institute of Physics. [doi:10.1063/1.3300129]

I. INTRODUCTION

The interest in the bonding between transition metal atoms and benzene began with the observation that certain colored solutions obtained by the addition of metal salts to liquid benzene arise from metal-benzene complexes. The interest on these complexes grew up soon after species, such as dibenzene chromium (Cr2C6H6)2 (Ref. 3), were synthesized and found to be very stable. In recent years, the investigation of sandwiches and other complexes of metal atoms has progressed because of their relevance to catalysis, magnetic and optical materials, polymers, spintronic devices, molecular sensors, and other applications (see, e.g., Refs. 4–17).

Despite of the difficulty of synthesizing these materials, the introduction of laser vaporization techniques in pulsed nozzle supersonic expansion sources, combined with mass spectrometry detection,18 has made possible studies of their bonding energetics, spectroscopic properties, and structures. Kurikawa et al.19 detected clusters containing transition metal atoms (M) and benzene molecules (Bz) with strong preference for species with [MnBzn+1]5+ stoichiometries. The neutral analogs have been recently studied by magnetic deflection.8 The stoichiometries of these clusters led to the proposal of a structure of sandwiches with the M atoms intercalated between benzene units. The magnetism observed in some of them was associated to the spin-coupling along the axis of the sandwich. Calculations for small MnBzn+1 clusters (n = 6) and for infinite (MBz)n+1 chains confirm the linear sandwich structure of these complexes as the most stable one.20 Recently, a theoretical study of the electronic and magnetic properties for (MBz)n polymers with M=Sc, Ti, V, Cr, and Mn has shown that all chains, except (CrBz)n, are metallic.21

Optical and UV spectroscopies are powerful tools for obtaining information on the electronic and atomic structure of clusters. Since the structure of a cluster is generally not directly accessible experimentally,22 its characterization can be done with a combination of optical spectroscopy and theoretical calculations.23 Optical absorption spectroscopy experiments for gas phase clusters containing transition metal atoms are quite challenging. Experiments and calculations of direct absorption or laser induced fluorescence are needed in the case of MnBzn+1 clusters. The possibility of varying the number of layers in the sandwich and the nature of the intercalated atoms raises expectations of interesting optical properties and possible applications of these materials.

We present first-principles calculations of the photoabsorption cross sections of MnBzn+1 sandwiches with M=Ti, V, Cr, and n ≤ 3, and for infinite (TiBz)n and (CrBz)n chains. The choice of Ti, V, and Cr is based on: (i) the experimental evidence that the tendency to form multidecker sandwiches is stronger for the early than for the late transition metals;19 (ii) other metal atoms, like silver or copper, have virtually no...
II. CALCULATION OF THE PHOTOABSORPTION SPECTRUM

A method extensively used to calculate the photoabsorption spectrum of clusters is the time-dependent density functional theory (TDDFT).23,25 The implementations used in practice have some limitations due to the approximate description of exchange and correlation effects between the electrons, like the known underestimation of the magnitude of the energy gap in semiconductors. A similar error could be expected for the gap between occupied and unoccupied states (HOMO-LUMO gap) in finite clusters. Possible errors in the gap and in the asymptotic behavior of the exchange-correlation (XC) potential in a finite cluster could affect the energies of the excited electronic states and thus the photoabsorption spectrum. For this reason, the TDDFT results are compared here with accurate results obtained from many-body perturbation theory.

We have first performed full optimizations of the geometries of these clusters, starting with the structures proposed in the literature and considering different spin-multiplicity states. DFT, implemented in the OCTOPUS package,26 was used with the PBE0 hybrid XC functional,27 which mixes a generalized gradient (GGA) functional with a predetermined amount (25 per cent) of exact exchange. The PBE0 functional improves the description of band gaps compared to uncorrected GGA functionals, through the incorporation of a fraction of Hartree–Fock exchange,28,29 and this provides a better starting point for TDDFT calculations. Semicore corrected norm-conserving pseudopotentials30,31 were employed. In agreement with previous work,9–11,19,20 our calculations show that the most stable structures of these complexes are sandwiches with the metal atoms intercalated and positioned in the axis of the sandwich (z-axis), as seen in Fig. 1. As previously reported for the V, Nb, and Ta systems,10,20 small rotation angles (1–3°) of the benzene rings relative to each other around the z-axis are found depending on the spin state. In the benzene molecules at the two ends of the sandwich, the planes of C and H hexagons are slightly shifted relative to each other along the z-axis by ∼0.05–0.10 Å. The infinite (MBz)n structures have been modeled as having pure Dnsh symmetry. The ground states of the TiBz+n structures and CrBz+n structures are nonmagnetic. The V2Bz3 clusters are more complex. The configuration with spin magnetic moment μz=0μB in which the two V atoms are antiferromagnetically coupled is very close in energy to a ferromagnetic configuration with μz=2μB in V2Bz3. The truly nonspin-polarized state is substantially above in energy (0.35 eV) with respect to the other two. In the case of V2Bz3, the magnetic configurations with μz=1μB and μz=3μB have very similar energies. For VBz2, the μz=1μB ground-state is clearly preferred.

Then, the photoabsorption spectrum is obtained using the TDDFT with the formalism developed by Casida.12 The ingredients for this calculation are the electronic single-particle energies and orbitals of the ground state described above and a XC kernel, for which the self-interaction corrected (SIC) adiabatic local density approximation (ALDA)33 is employed. The spectra of MxBz+n sandwiches with n=3 are shown in Fig. 1 for energies up to 6 eV. In the three series (M=Ti, V, Cr) the absorption energy thresholds decrease as more MBz units are added to the structure: the absorption threshold occurs near 4.5, 3.5, and 2.5 eV for the growing TiBz+n sandwiches, and near 3.0, 2.0, and 1.5 eV for the growing CrBz+n systems. In the more complex V3Bz4 structures, the red shift is also observed. However, two different spin states are nearly degenerate for V2Bz3 and the same occurs for V3Bz4, and the absorption threshold and the form of the spectrum are very sensitive to the spin state. It is likely that an experimental spectrum will be a superposition of the two spectra for each of those clusters. A general property is that the absorption intensity becomes higher as the number of layers in the sandwich increases.

The spectra of the TiBz+n sandwiches are characterized by a main peak centered at 4.7, 3.9, and 3.2 eV for n=1, 2, and 3, respectively. This peak corresponds to transitions from 3d states of Ti to benzene states. The intensity of the peak scales with the number of TiBz units forming the cluster. Two absorption regions can be distinguished in the spectra of CrBz2 and CrBz3: a low energy peak centered at 3.3 and 2.6 eV, respectively, followed by a broader absorption region at higher energies. Both spectra are similar except for a relative shift of about 0.6 eV. The origin of the low energy peaks is the same as in the clusters with Ti. The high energy region of the spectra arises from internal transitions in the metal atoms, from 3p states to the open 3d shell. The spectrum of

FIG. 1. Calculated TDDFT photoabsorption cross sections (in arbitrary units) for MxBz+n sandwich clusters, with n=3 and M=Ti, V, and Cr, for excitation energies up to 6 eV. Spectra for two nearly degenerate states with different spin magnetic moments are shown for V2Bz3 and V3Bz4. The excitation peaks have been broadened by Lorentzian functions. The geometrical structures are included as insets.
Cr$_3$Bz$_4$ differs from the other two. The overlap between the atomic 3d orbitals along the axis of the cluster lowers the energy of the atomic transitions and the second peak approaches closely to the first one. The spectra of V$_n$Bz$_{n+1}$ show some common features with those of the clusters containing Ti and Cr, that is, a shift of the spectrum to lower energies as $n$ increases. Of course, the analysis becomes more complex because different spin configurations appear for different $n$. An extended low-energy absorption region between 0 and 2 eV appears for the state of V$_3$Bz$_4$ with $\mu_z = 3\mu_B$. This arises from the increased occupation of orbitals of the spin up channel, giving rise to low energy transitions. The 3p–3d transitions discussed above for the Cr sandwiches occur at energies above 6 eV in the Ti and V sandwiches. The same 3p–3d transitions occur in the isolated Ti, V and Cr atoms at energies about 4.5–5 eV.

To explore the influence of quasiparticle and electron-hole interaction effects in the optical spectra of the sandwiches we have used the GW (Green function–screened interaction) approach and solved the Bethe–Salpeter equation (BSE), respectively. For this purpose: (i) as a starting point, taking the optimized geometries, the ground state electronic structure of Ti$_n$Bz$_{n+1}$ and Cr$_n$Bz$_{n+1}$ was obtained with the PWSCF code using an LDA XC–functional and norm-conserving pseudopotentials; (ii) electronic quasiparticle corrections were calculated within the GW approximation to the XC self-energy; (iii) finally, the BSE was solved for coupled electron-hole excitations, thereby accounting for the screened electron-hole attraction and the unscreened electron-hole exchange. Steps (ii) and (iii) were performed with the YAMBO code. In addition, infinite (TiBz)$_n$ and (CrBz)$_n$ chains have also been studied.

This formalism includes excitonic effects, which are more important in strongly correlated systems, and is able to distinguish the response to each individual light polarization component. In Fig. 2, a comparison between photoabsorption spectra calculated by the TDDFT and GW-BSE formalisms is shown. The GW-BSE spectra include separately the polarization contributions and the averaged spectra. For TiBz$_2$ and Ti$_2$Bz$_3$, the spectra obtained by both formalisms show some common features with those calculated with the local PZ parametrization. This is mainly due to the improved description of the HOMO–LUMO gap given by a GGA functional over the LDA. Although combining a GGA ground-state description with an LDA kernel is apparently not entirely consistent, this is comprised

FIG. 2. Calculated GW-BSE photoabsorption cross sections (in arbitrary units) of Ti$_n$Bz$_{n+1}$ and Cr$_n$Bz$_{n+1}$ sandwich clusters ($n=3$) and infinite chains. The individual x, y, and z polarization contributions and the spectrum averaged over the three polarizations are shown in different colors. TDDFT results are included for comparison. The inset in the bottom-left panel shows the GW energy band diagram for the infinite TiBz chain: $d_{xy}$, $d_{xz}$, $d_{yz}$ band (solid line); $d_{xy}$ band (dashed line); $d_{x^2−y^2}$, $d_{yz}$ band (dotted line).

excitation energies are not present in the GW-BSE spectra shown in Fig. 2. The reason is that those excitations are shifted to even higher energies. The pronounced size-dependent behavior of the optical spectrum, in which the visible absorption frequencies decrease with cluster size and saturate for the chain may possibly be used in tailoring the optical properties.

In Fig. 3, the TDDFT photoabsorption cross sections for TiBz$_2$ and CrBz$_2$, calculated by using two different XC functionals, the LDA of Perdew and Zunger (PZ) and the GGA-PBE0, with the same LDA+SIC kernel, are compared to the GW-BSE cross sections. The agreement with the GW-BSE spectrum is better for the TDDFT results obtained using the PBE0 hybrid XC functional, showing the main excitation peaks much closer to the GW-BSE profile than those calculated with the local PZ parametrization. This is mainly due to the improved description of the HOMO–LUMO gap given by a GGA functional over the LDA. Although combining a GGA ground-state description with an LDA kernel is apparently not entirely consistent, this is comprised.

FIG. 3. Comparison between the TDDFT photoabsorption cross sections for the TiBz$_2$ and CrBz$_2$ sandwich clusters, calculated by using two different XC functionals (LDA-PZ and GGA-PBE0), and the GW-BSE cross sections. The excitation peaks have been broadened by Lorentzian functions. The geometrical structures are shown as insets in Fig. 1.
of the doublet show a stronger overlap with the benzene orbitals (see Fig. 4). A salient feature of the infinite chain is that the \(d_{3z^2-r^2}\) band crosses the \(d_{xy}, d_{x^2-y^2}\) band (see inset in Fig. 2). In the (CrBz)\(_{\infty}\) chain, both the \(d_{3z^2-r^2}\) and the \(d_{xy}, d_{x^2-y^2}\) bands are filled, and the dispersion only gives rise to an intense redshift of the low energy absorption peak with respect to the CrBz\(_2\) spectrum (see Fig. 2). In contrast, the \(d_{3z^2-r^2}\) orbital is empty in TiBz\(_2\) and thus the dispersion leads to the metallization of the (TiBz)\(_{\infty}\). This fact has been already reported in a recent work by Xiang et al.,\(^{21}\) who used the PW91 functional\(^{40}\) in their work. However, it is well known that DFT tends to underestimate the gaps, predicting a metallic behavior for small gap semiconductors. This trend is corrected by the GW calculations,\(^{41}\) so we propose the metallization of the (TiBz)\(_{\infty}\) chain predicted by Xiang et al. as a reliable effect, with possible interest in applications involving tunable electro-optical properties in nanoscale devices. The present results also predict that the (CrBz)\(_{\infty}\) chain does not become metallic, again in agreement with Xiang et al.\(^{21}\)

It is worthwhile to apply a similar orbital-based analysis to V\(_3\)Bz\(_4\) in its two different spin configurations, in order to shed light on the source of the differences seen in the low-energy region of the spectra for the magnetic states with \(\mu = 1 \mu_B\) and \(\mu = 3 \mu_B\). The ordering of the electronic levels and the form of the associated orbitals are depicted in Fig. 5. The low energy peaks in the photoabsorption spectrum of the state with \(\mu = 3 \mu_B\) arise from excitations from its HOMO orbital. This molecular orbital corresponds to an antisymmetric linear combination of the \(d_{3z^2-r^2}\) orbitals of the two V atoms located at the extremes of the sandwich. It is remarkable that this is the only one of the last four frontier orbitals with ungerade symmetry. Low energy transitions are allowed from this level, but not from the HOMO-1, HOMO-2, or HOMO-3 levels. On the other hand, that orbital is empty in the case of \(\mu = 1 \mu_B\); therefore, there will be no transitions from this orbital when \(\mu = 1 \mu_B\).

Some photoemission experiments have been performed

**FIG. 4.** Frontier orbitals of TiBz\(_2\) (top panel) and CrBz\(_2\) (bottom panel), and axis set chosen in the calculations. On the right hand side, the orbitals are labeled in accordance with the D\(_{6h}\) point group symmetry. The thick vertical red arrows indicate the transitions responsible for the low energy peaks of Fig. 2.

**FIG. 5.** Frontier orbitals of V\(_3\)Bz\(_4\) for its two different spin configurations with \(\mu = 1 \mu_B\) and \(\mu = 3 \mu_B\). On the left and right hand sides, the orbitals are labeled in accordance with the D\(_{6h}\) point group symmetry.
on the neutral and anionic species of several of these sandwiches. Recently, Sohnlein et al. have performed ZEKE-PES measurements on bis-benzene complexes of titanium and vanadium, which provide information on the energetics of the occupied orbitals, but not of the unoccupied ones, and hence they cannot be compared with the optical spectra. On the other hand, Zheng et al. have performed anion photoelectron measurements on \((\text{Ti}_n\text{Bz}_m)\). These experiments provide information on the energy of the LUMO of the neutral \(\text{Ti}_n\text{Bz}_m\) system with respect to its occupied orbitals. It can be inferred from that work that the LUMO of TiBz$_2$ is about 1 eV above the occupied states, in good agreement with our calculations in which the HOMO-LUMO gap is 1.13 eV. However, this energy is much lower than the one obtained for the first intense peak in the photoabsorption spectrum, namely 4 eV in the GW-BSE calculations. This is due to the fact that all the transitions from the occupied frontier orbitals of TiBz$_2$ to its LUMO are forbidden by symmetry. The same feature occurs for the rest of the \(\text{Ti}_n\text{Bz}_{n+1}\) systems studied in this work.

### III. CONCLUSION

A suitable combination of TDDFT and GW-BSE calculations has allowed an exhaustive analysis of the optical spectra of sandwich clusters formed by transition metal atoms intercalated between parallel benzene molecules. The low excitation energy peaks are characteristic of the robust bonding in this kind of complexes, and different metal atoms and cluster sizes could be used to tune the optical properties according to specific functionality targets. The differences in the spectra can be used to identify relative abundances of isomers with different spins in experimental studies. The present results are in good agreement with the experiments and calculations found in recent literature.

### ACKNOWLEDGMENTS

This work was supported by MICINN of Spain (Grant No. MAT2008-06483-C03-01) and Junta de Castilla y León (Grant Nos. VA017A08 and GR23). J.A.A. acknowledges an Ikerbasque fellowship from the Basque Foundation for Science, and the hospitality of DIPC. J.M.G.-L. acknowledges funding from Spanish MICINN through Juan de la Cierva and José Castillejo programs.