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Kyoto University
Critical reinvestigation of vibronic couplings in picene from view of vibronic coupling density analysis

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Vibronic coupling constants in the monoanionic, trianionic, and excited states of picene are evaluated from the total energy gradients using the density functional theory. Employing the calculated vibronic coupling constants in the excited state of a neutral molecule, an electron energy loss spectrum (EELS) is simulated to be compared with the experimental spectrum. The calculated vibronic coupling constants are analyzed in terms of the vibronic coupling density which enables us to analyze vibronic couplings based on the relation between the electronic and vibrational structures. The vibronic coupling constants reported by Kato et al. [J. Chem. Phys. 116, 3420 (2002); Phys. Rev. Lett. 107, 077001 (2011)] are critically discussed based on the vibronic coupling density analysis.

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After the discovery of superconductivity in alkali-metal (K, Rb) doped picene,1 experimental and theoretical studies on the electronic structure on the picene have accumulated.2–14 The vibronic coupling (electron-vibration coupling)15 is an important interaction in electronic properties such as superconductivity. Okazaki et al. have discussed the importance of the vibronic couplings in doped picene based on their photoelectron spectra (PES).2 Therefore, evaluation of the vibronic coupling constants (VCCs) is crucial to discuss the vibronic progressions in the EELS of the intramolecular electronic and vibrational structures. On the basis of the VCD analysis, we have succeeded in designing carrier-transporting molecules with small vibronic couplings, which are required for the electronic and vibrational structures. The vibronic coupling constants reported by Kato et al. [J. Chem. Phys. 116, 3420 (2002); Phys. Rev. Lett. 107, 077001 (2011)] are critically discussed based on the vibronic coupling density analysis.

We evaluated VCCs of mode \( \alpha \), \( V_\alpha \), from the gradients of the total energies using the density functional theory. Employing the calculated vibronic coupling constants in the excited state of a neutral molecule, an electron energy loss spectrum (EELS) is simulated to be compared with the experimental spectrum. The calculated vibronic coupling constants are analyzed in terms of the vibronic coupling density which enables us to analyze vibronic couplings based on the relation between the electronic and vibrational structures. The vibronic coupling constants reported by Kato et al. [J. Chem. Phys. 116, 3420 (2002); Phys. Rev. Lett. 107, 077001 (2011)] are critically discussed based on the vibronic coupling density analysis.

\[
V_\alpha = \langle \Psi | \left( \frac{\partial \hat{H}}{\partial Q_\alpha} \right)_{R_0} | \Psi \rangle = \left( \frac{\partial E}{\partial Q_\alpha} \right)_{R_0},
\]

where \( \hat{H} \) denotes a molecular Hamiltonian, \( R_0 \) is the equilibrium geometry of the ground state of the neutral picene, \( \Psi \) is an electronic wave function of the excited or anionic state at \( R_0 \), and the phase of a normal mode \( \alpha \) is chosen so that \( V_\alpha \) becomes negative. The vibronic Hamiltonian is written as

\[
\hat{H}_{\text{vibro}} = \sum_\alpha \left[ \hat{T}(Q_\alpha) + \frac{1}{2} \omega_\alpha^2 Q_\alpha^2 + V_\alpha Q_\alpha \right],
\]

where \( \hat{T}(Q_\alpha) \) denotes the kinetic energy operator of a vibration and \( \omega_\alpha \) is the vibrational frequency. We employed Becke’s hybrid functional (B3LYP)25 and Perdew and Wang’s one with a generalized gradient approximation (PW91)26 with the triple-zeta 6-311 + G(d,p) basis set. The geometries were optimized for the neutral ground state. The optimized structures with \( C_{2v} \) symmetry were checked with a vibrational analysis to be a minimum. The time-dependent density functional theory is applied for the excited-state calculations. We performed analytical force calculations for the excited and the anionic states to obtain the VCCs.27 The electronic and vibrational structures as well as the forces were obtained using a program package, GAUSSIAN 09.27 The VCCs were calculated using our codes.

Since the fine structure observed in the EELS by Roth et al. can originate from the vibronic couplings which they did not take into account in Ref. 3, we simulated the EELS to find an appropriate functional for the VCC calculations. EELS was simulated employing the same method as described in Ref. 18. We considered a thermal excitation at 20 K, where Roth et al. observed the EELS. The calculated excitation energies are 3.7040 and 3.2520 eV for the B3LYP and PW91 functionals, respectively. The result using the PW91 functional reproduces well the experimental excitation energy 3.25 eV.

The calculated VCCs in the \( \Sigma_2 \) state, we simulated the EELS (Fig. 1). In the simulations, the 0-0 transition is set to 3.24 eV, and the linewidth \( \sigma \) is assumed to be 270 cm\(^{-1}\) [the full width at half maximum (FWHM) is 39.4 meV].
The calculated VCCs$^{28}$ as well as the vibrational frequencies are tabulated in the Supplemental Material.$^{29}$ The spectrum calculated employing the PW91 functional [Fig. 1(a)] shows a better fit than that using the B3LYP functional [Fig. 1(b)]. In the calculation employing the B3LYP, the second strongest vibronic couplings around 1350 cm$^{-1}$ are estimated to be larger than those in the result using the PW91.$^{30}$ Therefore, the relative intensities of EELS in the lower energy are reproduced employing the PW91.$^{31}$ Judging from the calculations of the excitation energies and the simulated spectra, hereafter we employ the PW91 functional for the calculation of the ionic states. Further comparisons of functionals are out of the scope of the present Rapid Communication.

We calculated the VCCs $V_a$ in the monoanion ($^1A_2$), dianion ($^1A_1$), and trianion ($^2B_3$) (see the Supplemental Material$^{29}$). Kato et al. have calculated orbital vibronic coupling constants (OVCCs) of the lowest unoccupied molecular orbitals (LUMOs) and next LUMOs (NLUMOs)$^{11,16,17}$ Hence we also calculated the OVCCs $V_{i,\alpha}$ from the VCCs $V_a$ for comparison with the ones previously reported. The vibronic Hamiltonian (2) is mapped onto a model Hamiltonian. The model Hamiltonian considered in Refs. 10 and 11 is written as follows:

$$\hat{H}_{\text{vibro}} = \sum_{\alpha,i,\sigma} \hbar \omega_{\alpha} \left[ \hat{b}_{\alpha}^\dagger \hat{b}_{\alpha} + \frac{g_{i,\alpha}}{\sqrt{2}} (\hat{b}_{\alpha}^\dagger + \hat{b}_{\alpha}) \hat{c}_{i\sigma}^\dagger \hat{c}_{i\sigma} \right],$$  (3)

where orbitals $i$ are LUMO (L) and NLUMO (NL), and $\alpha$ runs over all the active $a_1$ modes.$^{28}$ The dimensionless OVCC $g_{i,\alpha}$ is defined by $g_{i,\alpha} = V_{i,\alpha} / \sqrt{\hbar \omega_{\alpha}}$. $\hat{c}_{i\sigma}^\dagger$ ($\hat{c}_{i\sigma}$) is the creation (annihilation) operator of orbital $i$ and spin $\sigma$, and $\hat{b}_{\alpha}^\dagger$ ($\hat{b}_{\alpha}$) is the creation (annihilation) operator of mode $\alpha$. The OVCCs of LUMO $V_{L,\alpha}$ and NLUMO $V_{NL,\alpha}$ are obtained from the VCCs of monoanion $V_{\text{mono},\alpha}$ and the difference between the

![Graph](image1)

**FIG. 1.** (Color online) Red lines are simulated electron energy loss spectra (EELS) calculated by (a) the PW91 and (b) the B3LYP functional. Blue dots indicate the experimental one by Roth et al.$^5$ The 0-0 transition is set to 3.24 eV. The other parameters employed in the simulations are $T = 20$ K and $\sigma = 270$ cm$^{-1}$.

![Graph](image2)

**FIG. 2.** (Color online) Calculated (top) orbital vibronic coupling constants $V_{i,\alpha}$ and (bottom) electron-phonon couplings $\lambda_{i,\alpha}/N(0)$ in the (a) monoanion and the (b) trianion. The subscripts L and NL denote LUMO and NLUMO, respectively. The insets show the orbital vibronic couplings between 1300 and 1650 cm$^{-1}$.
indicate positive and negative, respectively. The threshold is 1.0 \times 10^{-2} \text{ a.u.}

VCCs of trianion \( V_{\text{tri},\alpha} \) and dianion \( V_{\text{di},\alpha} \), respectively: \( V_{\text{mono},\alpha} = V_{\text{mono}} - V_{\text{di},\alpha} \). It should be noted that the present OVCCs effectively incorporate the contributions from all the occupied orbitals, which are important in quantitative arguments.\(^{18}\) The OVCCs \( V_{\alpha} \) and intramolecular electron-phonon couplings \( \lambda_{\alpha}/N(0) = \sum_{\alpha} V_{\alpha}^2/\omega_{\alpha}^2 \) are shown in Fig. 2 and are tabulated in the Supplemental Material.\(^{29}\) \( N(0) \) is the density of states at the Fermi level.

In both the monoanion and trianion, two sets of modes with strong VCCs are observed: (1) the maximal coupling modes around 1600 cm\(^{-1}\) and (2) the second group of strong modes around 1350 cm\(^{-1}\). This is qualitatively consistent with the calculations by Subedi and Boeri.\(^{10,32}\) However, in the calculations of Refs. 11, 16, and 17,\(^{33}\) the modes around 1600 cm\(^{-1}\) are weak for both the monoanion and trianion. The total electron-phonon couplings \( \lambda_{L}/N(0) = \sum \lambda_{L,\alpha}/N(0) \) and \( \lambda_{\text{SL}}/N(0) = \sum \lambda_{\text{SL},\alpha}/N(0) \) are 196.6 and 119.9 meV, respectively. The total coupling of the trianion \( \lambda_{\text{SL}}/N(0) \) is in line with that of Subedi and Boeri (110 \pm 5 meV).\(^{10}\) On the other hand, in Ref. 11, \( \lambda_{L}/N(0) = 178 \text{ meV} \) and \( \lambda_{\text{SL}}/N(0) = 206 \text{ meV} \). They underestimated and overestimated the total couplings for the monoanion and trianion, respectively.

We will discuss the disagreement from the view of the electronic and vibrational structures. The calculated VCCs can be rationalized based on the vibronic coupling density.\(^{20,21}\) A VCD \( \eta_\alpha \) is defined as

\[
\eta_\alpha(\mathbf{r}) = \Delta \rho(\mathbf{r}) \times v_\alpha(\mathbf{r}),
\]

where \( \Delta \rho(\mathbf{r}) = \rho(\mathbf{r}) - \rho_0(\mathbf{r}) \) is the electron density difference between the electron density of an ionic state \( \rho \) and that of a neutral state \( \rho_0 \). The potential derivative \( v_\alpha(\mathbf{r}) \) is the derivative with respect to a mass-weighted normal coordinate \( Q_\alpha \) of the potential \( u(\mathbf{r}) \) acting on one electron at a position \( \mathbf{r} \) from all the nuclei. The vibronic coupling constant is equal to the integral of \( \eta_\alpha(\mathbf{r}) \) over space \( \mathbf{r} \):

\[
V_\alpha = \int d^3 \mathbf{r} \eta_\alpha(\mathbf{r}). \tag{5}
\]

The VCD gives a local picture of the vibronic coupling, and hence enables us to discuss the strength of the coupling qualitatively.

We will concentrate on the \( a_1(27) \) mode of the maximal-coupling mode around 1600 cm\(^{-1}\) and the \( a_1(21) \) mode from the second group around 1350 cm\(^{-1}\) because the \( a_1(27) \) and \( a_1(21) \) modes are close to mode 5 in Ref. 10 and the 21st mode in Ref. 11, respectively. Figure 3 shows the potential derivatives \( v_\alpha(\mathbf{r}) \) for the \( a_1(27) \) and \( a_1(21) \) modes. The distribution of the potential derivative with respect to the \( a_1(27) \) mode is located on the armchair edges of the central three hexagons. On the other hand, that with respect to the \( a_1(21) \) mode is on the terminal hexagons.

Figure 4(b) shows the electron density difference \( \Delta \rho \) of the monoanion. Since an additional electron occupies the LUMO [Fig. 4(a)], the positive \( \pi \) density (white) appears in \( \Delta \rho \). It should be noted that there occurs a decrease of the \( \sigma \) density (blue) in the molecular plane. Such a polarized density originates from the Coulomb interactions between the electrons occupying the LUMO and all electrons in the doubly occupied orbitals below the highest occupied molecular orbital (HOMO).\(^{20}\)

Figure 5 shows the vibronic coupling densities \( \eta_\alpha(\mathbf{r}) \) of the monoanion with respect to the \( a_1(27) \) and \( a_1(21) \) modes. As for the \( a_1(27) \), the electron density difference \( \Delta \rho(\mathbf{r}) \) shows a considerable overlap with the potential derivative \( v_\alpha(\mathbf{r}) \). On
the other hand, $\Delta \rho(\mathbf{r})$ does not significantly overlap $v_\alpha(\mathbf{r})$ for the $a_1(21)$ mode. Therefore, the VCD $\eta_\alpha$ (4) for the $a_1(27)$ is larger than that for the $a_1(21)$. Particularly, $\eta_\alpha$ for the $a_1(27)$ has a large distribution on the central three armchair edges. Accordingly, the VCC of the $a_1(27)$ mode is larger than that of the $a_1(21)$ mode.

The VCD $\eta_\alpha$ for the $a_1(27)$ mode on the bonds appears due to the polarization of $\Delta \rho$, thus neglecting such a polarization can give rise to qualitative, or sometimes quantitative, errors in VCC calculations based on the orbital levels. In addition, an electron density difference is usually different from the orbital density of HOMO or LUMO. The many-body effect sometimes plays a crucial role on the electron density difference, and therefore vibronic couplings.

Kato et al. have calculated VCCs as the gradients of the orbital energies (orbital gradients) with respect to normal coordinates. We also obtained the orbital gradients of selected modes from the neutral LUMO $\partial \epsilon_L/\partial Q_\alpha$ and from the singly occupied molecular orbital (SOMO) of the monoanion $\partial \epsilon_S/\partial Q_\alpha$ to compare the present VCCs calculated analytically from the total energy gradient as well as numerical gradients. The orbital gradients and the numerical energy gradients were obtained by fitting linear and quadratic polynomials, respectively, in the range from $-0.2$ to $0.2$ a.u. with a step size $0.05$ a.u. where $a_0$ is the Bohr radius. The range and the step size could be different from those of Kato et al. We summarize the results of the calculations in Table I. All the results indicate that the vibrational modes around $1600$ cm$^{-1}$ have the maximal coupling.

![FIG. 5. (Color online) Vibronic coupling densities for (a) $a_1(27)$ ($1605$ cm$^{-1}$) and (b) $a_1(21)$ ($1379$ cm$^{-1}$) modes of the monoanion. The white and blue areas indicate positive and negative, respectively. The threshold is $2.0 \times 10^{-3}$ a.u.](image)

Though the gradients of the LUMO level can yield results which are qualitatively consistent with the gradients of the total energy, the results in Refs. 11 and 16 are not the case. In their calculations, the vibrational mode around $1380$ cm$^{-1}$ has the maximal coupling, $2.3 \times 10^{-4}$ a.u. for the monoanion. However, since $\Delta \rho$ [see Fig. 4(b)] is mainly located on the armchair edges of the central three hexagons, it does not overlap with the potential derivatives $v_\alpha$ of the mode [Fig. 3(b)], the VCC of the mode cannot be the maximal. A similar discussion holds for the mode around $1520$ cm$^{-1}$ which has a maximal coupling of $2.6 \times 10^{-4}$ a.u. for the trianion.

In summary, we calculated the VCCs of picene for the excited state $^1B_2$ of the neutral molecule, the monoanionic, and the trianionic states of the molecule from the gradients of the total energies with respect to the normal modes. In previous studies, the VCCs calculated form the gradients of the orbital energies of the frontier level, LUMO. In other words, they have regarded the OVCCs of the LUMO as the VCCs. The present approach can provide quantitatively reliable VCCs, since all the occupied orbitals can contribute to the VCCs due to the selection rule. This has been discussed in Ref. 18 in detail. We simulated the EELS and compared the spectra with the experimental spectrum by Roth et al. Needless to say, a comparison between the theoretical and the experimental results on the vibronic couplings in picene is necessary to obtain reliable VCCs. From these simulations, we determined an appropriate functional for the calculation of the vibronic couplings in picene. Using the functional which reproduces the EELS, we evaluated the vibronic coupling constants of the picene anions. The calculated vibronic couplings of the anions can be employed for assignments of spectra, for example, photoelectron spectra of the anions. We discussed the vibronic couplings of the picene anions in terms of the vibronic coupling density. Based on the analysis, the present vibronic couplings are reasonable compared with the values reported previously.

Numerical calculations were performed partly in the Supercomputer Laboratory of Kyoto University and Research Center for Computational Science, Okazaki, Japan. This work was supported in part by the Japan Society for the Promotion of Science (JSPS) through its Funding Program for the Global COE Program “International Center for Integrated Research and Advanced Education in Materials Science” (No. B-09) of the Ministry of Education, Culture, Sports, Science and Technology (MEXT) of Japan.
From the selection rule for the vibronic coupling, the excited and anionic electronic states couple only to the $a_1$ modes.

In terms of the electron-phonon couplings, the total coupling in the excited state obtained by the B3LYP (355.8 meV) is larger than that by the PW91 (291.4 meV) by 64 meV.

The peaks in the higher energy than 3.6 eV include other excitations (Ref. 5), while we do not consider them in our simulation. Thus the simulated spectrum does not agree with the experimental one in the energy range. Since we focus on $S_2$, we do not consider the higher excitations.

Subedi and Boeri calculated the electron-phonon couplings using the density functional perturbation theory (Ref. 34) within the local density approximation.

Kato et al. calculated the electron-phonon couplings from the gradients of orbital level using the B3LYP functional.