Enhancement of parity-violating energy difference of CHFClBr, CHFClI, and CHFBrI by breaking the cancellation among valence orbital contributions

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The enhancement of the parity-violating energy difference (PVED) by electronic excitation is studied for H₂X₂ (X = O, S, Se, Te), CHFClBr, CHFClI, and CHFBrI. To clarify the enhancement mechanism, the dihedral angle dependence of the PVED of H₂X₂ in excited states is studied. If the contribution from the highest occupied molecular orbital (HOMO) to the PVED in the ground state is larger than the sum of those from all occupied orbitals, the PVED in the first excited state has a much larger value compared to the ground state due to cancellation breaking among valence orbital contributions. This enhancement is named cancellation breaking enhancement. The PVED enhancement is also studied for CHFClBr, CHFClI, and CHFBrI in excited states, and the cancellation breaking enhancement is confirmed. When the PVED contribution from the HOMO is larger than any other contribution, the cancellation breaking enhancement hypothesis provides the estimate of PVED in the first excited state from the HOMO contribution.

I. INTRODUCTION

Parity-violating energy difference (PVED) is the energy difference between an enantiomeric pair, which is induced by the weak interaction between nuclei and electrons in a molecule. PVED is studied by many groups in experimental and theoretical fields and is one subject related to the origin of homochirality in nature [1]. In addition, if PVED is experimentally observed, it is the first confirmation of the imprint of the weak interaction on molecules. A subsequent comparison of the measured PVED value with theoretical prediction may constrain new physics beyond the standard model in particle physics, since many extended models have new particles contributing to the PVED. Hence, PVED may provide a new tool wherein molecules are used to investigate new physics beyond the standard model.

Thus far, PVED has not been experimentally observed, because of its small size. The tightest upper bound comes from the experiment using the CHFClBr molecule [2]. In this experiment, the vibrational frequency difference between the enantiomeric pair of the molecule was measured and |Δν/ν| < 2.5 × 10⁻¹³ was reported. This bound constrains the PVED with the relation, E_{el}^{PV} / E_{el} ∼ E_{el}^{PV} / E_{vib} [3].

The usage of molecules with large PVED, in addition to the improvement of experimental technology, is important for the observation of nonzero PVED in future experiments. The PVED of chiral molecules is predicted by quantum chemistry computations with relativistic effects. Many works [4–6] have reported larger PVED values than CHFClBr, and we consider that the search for molecules with larger PVED is still important. It is known that molecules with heavy elements have large PVED values due to their large spin-orbit coupling, but computations of the PVED of these molecules require huge costs. It is impossible to exhaustively compute the PVED of molecules with heavy elements. Hence, we take another route to hunt for molecules with large PVED: usage of the electronic excitation of chiral molecules.

In Ref. [7], it was proposed that the PVED of electronic excited or ionic states of a chiral molecule is significantly enhanced compared to the neutral ground state. The contributions to the PVED from valence orbitals are canceled out in the ground states of many chiral molecules, and the PVED is expected to be enhanced significantly if this cancellation is broken by excitation or ionization. In our previous paper [8], this speculation was confirmed for H₂X₂ (X = O, S, Se, Te) by Coupled Cluster Singles and Doubles (CCSD) and Equation-of-Motion CCSD computations. In the work, it was reported that the PVED enhancement in the optimized H₂X₂ structure is much larger than the structure with φ = 45° where φ is the dihedral angle of H₂X₂. In the present paper, the PVED enhancement mechanism is studied in detail through the dependence on the dihedral angle. In addition, we confirm that this enhancement occurs in other molecules: CHFClBr, CHFClI, and CHFBrI. CHFClBr was used in the observational experiment [2]. CHFClI and CHFBrI are its derivatives that were studied as candidates for experimental improvement [5–9].

This paper is organized as follows. In the next section, the definition of the PVED is introduced briefly. Then, our computational method and details are described in Sec. [III] In Sec. [IV] our results are presented. The dependence of the PVED on the dihedral angle of H₂X₂ molecules is studied and one mechanism of PVED enhancement is proposed. Then, PVED enhancement by electronic excitation is demonstrated for CHFClBr, CHF-
CII, and CHFBrI, and our hypothesis of the enhancement mechanism is confirmed for these molecules. The last section is devoted to our conclusion and discussion.

II. THEORY

The PVED is defined as twice the parity-violating energy, $E_{PV}$, 

$$\Delta E_{PV} = 2|E_{PV}|.$$  

(1)

The dominant part of the PVED is calculated by the following equation,

$$E_{PV} = \frac{G_F}{2\sqrt{2}} \sum_n g_\nu^0 M^n_{PV},$$  

(2)

where $G_F = 1.663788(6) \times 10^{-5} (\hbar e)^3 \text{ GeV}^{-2}$ is the Fermi coupling constant [10]. The coupling of the nucleus, $n$, is given as $g_\nu^0 = Z^n(1 - 4 \sin^2 \theta_W) - N^n$, where $Z^n$ and $N^n$ are the number of protons and neutrons in the nucleus $n$, respectively, and $\theta_W$ is the weak-mixing angle, $\sin^2 \theta_W = 0.23121(4)$ [10]. The definition of $M^n_{PV}$ is the electron chirality at the nucleus $n$,

$$M^n_{PV} = \int d^3x (\Psi^\dagger |\chi_{e,x}^i(x)\gamma_\nu\chi_{e,x}^i(x)|\chi_{n,x}^i(x)\Psi),$$  

(3)

where $\chi_{e,x}^i$ and $\chi_{n,x}^i$ are the field operators of the electron and the nucleus $n$, respectively, $|\Psi\rangle$ represents the state vector, and $\gamma_\nu$ is defined with gamma matrices, $\gamma_5 = \gamma^5 = i\gamma^0\gamma^1\gamma^2\gamma^3$. The ordinary Dirac representation [11] is adopted in this article. In the derivation of this equation, protons and neutrons are assumed to have the same distribution in nuclei. Since the distribution of nuclear density is strongly localized, $M^n_{PV}$ is almost equal to the electron chirality at the nucleus position, where the electron chirality density is defined as $\langle \Psi^\dagger |\chi_{e,x}^i(x)\gamma_\nu\chi_{e,x}^i(x)|\Psi \rangle$. The derivation of Eq. (3) is explained in detail in our previous paper [8].

In the current study, a contribution to $M^n_{PV}$ from the $i$-th orbital is used and is defined as

$$M^n_{PV,i} = \int d^3x (\Psi^\dagger |\chi_{e,x,i}^i(x)\gamma_\nu\chi_{e,x,i}^i(x)|\chi_{n,x}^i(x)\Psi),$$  

(4)

where $\chi_{e,x,i}$ corresponds to the electron in the $i$-th orbital. In this expression, two degrees of the Kramers pair are separately treated, and $M^n_{PV} = 2\sum_i M^n_{PV,i}$ in closed-shell molecules.

III. COMPUTATIONAL DETAIL

The expectation values of $E_{PV}$ in the excited states are calculated using the Finite-Field Perturbation Theory (FFPT) [12-14] with the EOM-CCSD method [15]. For confirmation of our FFPT computations, the values of $E_{PV}$ in the ground states of our target molecules are compared with values based on the Z-vector method in CCSD computations [16].

In our computations, $E_{PV}$ is calculated as the sum of contributions from nuclei, $E_{PV} = \sum_n E^n_{PV}$ where $E^n_{PV}$ is the contribution to $E_{PV}$ from the nucleus $n$. In the FFPT method, $E^n_{PV}$ is calculated by the perturbation method whose perturbation Hamiltonian is chosen to be $H^n_{PV} = \gamma^n \int d^3x \chi_{e,x}^i(x)\gamma_\nu \chi_{e,x}^i(x)\chi_{n,x}^i(x)$, where $\lambda^n$ is the perturbation parameter for the nucleus $n$, and $\langle \Psi |H^n_{PV}|\Psi \rangle = \lambda^n M^n_{PV}$. The Hellmann-Feynman theorem tells us the following relation:

$$\frac{\partial E^n(\lambda^n)}{\partial \lambda^n} \bigg|_{\lambda^n=0} = \left\langle \Psi \left| \frac{\partial H}{\partial \lambda} \right| \Psi \right\rangle = M^n_{PV},$$  

(5)

where $H$ is the total Hamiltonian. The computation of the derivative $\partial E^n(\lambda^n)/\partial \lambda^n|_{\lambda^n=0}$ in Eq. (5) is performed using the finite difference method as $\partial E^n(\lambda^n)/\partial \lambda^n|_{\lambda^n=0} = (E^n(\lambda^n) - E^n(-\lambda^n))/(2\lambda^n)$. For this computation, the choice of the value of $\lambda^n$ is important. For the accurate calculation of the derivative in the finite difference method, small $\lambda^n$ is suitable for avoiding contamination by the effect of second or higher order derivatives, while with too small $\lambda^n$, the effect of $H^n_{PV}$ is less than computational error. Hence, an appropriate value of $\lambda^n$ should be adopted. The determination of the $\lambda^n$ value is based on a comparison between the FFPT and Z-vector results in the ground state. For excited states, the same values of $\lambda^n$ are used. Additionally, we have confirmed that the values of $E_{PV}$ in excited states are independent of $\lambda^n$ around the appropriate value of $\lambda^n$. For the computation of $E_{PV}$, the contribution from hydrogen atoms is neglected, since the coupling $g_\nu^0 \approx 0.0752$ is much smaller than other atoms and $M^n_{PV}$ is small due to the smallness of the spin-orbit interaction.

Our targets for the $E_{PV}$ computations are $H_2X_2$ ($X = O$, S, Se, and Te), CHFClBr, CHFCII, and CHFBrI molecules. The structures of $H_2X_2$ are the same as those in our previous paper [8] except for the dihedral angle $\phi$, which is treated as a free parameter in the present work. The structures of CHFClBr, CHFCII, and CHFBrI are determined by geometrical optimization computations with GAUSSIAN16 [17]. Density functional theory (DFT) with Becke’s three-parameter Lee-Yang-Parr (B3LYP) [18] basis sets [19] with the Stuttgart-Dresden relativistic effective core potential (Br and I) [20] are adopted. The structures of excited states are chosen to be the same as the ground states for all molecules.

The evaluation of Eq. (3) requires relativistic four-component wave functions for electrons, and the DIRAC program is used for these electronic structure computations. For computations of $H_2X_2$ molecules, the DIRAC19 program [22] is used, while DIRAC21 [23] is used for CHFClBr, CHFCII, and CHFBrI. For the self-consistent field (SCF) method computations, the Dirac-Coulomb-Gaunt Hamiltonian is adopted, while molecular mean-field approximations [25] to the Dirac-Coulomb-Gaunt Hamiltonian is adopted.
Coulomb-Gaunt Hamiltonian are adopted for post-SCF computations, i.e., CCSD and EOM-CCSD computations. For the latter computations, the RELCCSD module is used [15, 26, 27].

In computations of H$_2$X$_2$ molecules, dyall.acv3z [28] is employed and this choice was confirmed to be sufficiently accurate in our previous paper [5]. In CCSD and EOM-CCSD computations of H$_2$O$_2$, all electrons are treated as correlating orbitals and all virtual orbitals in active space are taken into account. In computations of H$_2$S$_2$, H$_2$Se$_2$, and H$_2$Te$_2$, correlating orbitals are chosen to be 2s2p, 4s4p, and 5s5p, respectively, and the cutoff of virtual orbitals in the active space is 10 Hartree (H$_2$S$_2$ and H$_2$Se$_2$) and 70 Hartree (H$_2$Te$_2$). The convergence criterion for EOM-CCSD computations is chosen to be $10^{-10}$ as used in our previous work [5]. The distribution of the electron chirality density is calculated with the QEDynamics program package [29] for the wave function computed by the Hartree-Fock (HF) method of DIRAC19.

In the computations of CHFClBr, CHFClI, and CHFBrI molecules the dyall.acv2z basis set [28] is adopted, and the difference in $E_{PV}$ among basis sets is discussed in a later section. In the CCSD computations, the cutoff of virtual orbitals in the active space is 50 Hartree, and correlating orbitals are chosen to be 3d4s4p(Br), 3s3p(Cl), 2s2p(F), 2s2p(C), 1s(H), and 4d5s5p(I) for any molecules. In EOM-CCSD computations of these molecules, the convergence criterion is loosened compared to that of H$_2$X$_2$ to 10$^{-7}$ as used in our previous work [5]. The distribution of the electron chirality density is calculated with the QEDynamics program package [29] for the wave function computed by the Hartree-Fock (HF) method of DIRAC19.

In the following, the atomic units are employed and in particular $E_{PV}$ is expressed in Hartree ($E_h$).

A. H$_2$X$_2$

In our previous paper [5], the magnitude of the enhancement of $E_{PV}$ in H$_2$X$_2$ is different between the optimized and $\phi = 45^\circ$ structures. Hence, H$_2$X$_2$ is chosen as a model and the dependence of the enhancement on the dihedral angle is investigated. In this calculation, the perturbation parameter $\lambda^n$ in the FFPT method is set to $10^{-3}$, $10^{-3}$, $10^{-4}$, and $10^{-5}$ (a.u.), for O, S, Se, and Te, respectively, which are determined in our previous paper [5] by comparing $E_{PV}$ values in the ground state with those calculated by the Z-vector method in CCSD computations.

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In Fig. 1, the dependence of $E_{PV}$ on the dihedral angle is shown. The enhancement is strongly dependent on the dihedral angle and $E_{PV}$ of H$_2$X$_2$ exhibits some common peaks. For the excited states of 1a and 1b, there is a peak with positive $E_{PV}$ at $\phi = 90^\circ$. For the 2a state, the positive peak is found at $\phi = 120^\circ$ ($X=O$) and $105^\circ$ ($X=S$, Se, Te). For the 2b state of H$_2$O$_2$, a negative bump is seen around $\phi = 75^\circ$, while for H$_2$S$_2$, H$_2$Se$_2$, and H$_2$Te$_2$ the positive peak and negative bump are observed at $\phi = 75^\circ$ and $\phi = 90^\circ$, respectively. Since the optimized dihedral angle of H$_2$X$_2$ is $\phi = 115^\circ$ ($X=O$) and $\phi = 90^\circ$ ($X=S$, Se, Te), large enhancement occurs in the optimized structure. In this calculation of $E_{PV}$, the value of $\lambda^n$ is fixed for all values of $\phi$. This treatment is sufficiently accurate for our study. The deviation of $E_{PV}$ from the Z-vector result is within a few percent except for $\phi = 90^\circ$, where the deviation is 1-7% due to the smallness of $E_{PV}$ itself.

To understand this spectrum, a contribution from each orbital is studied. Figure 2 shows $M_{PV,i}^X$ of HOMO, HOMO-1, and HOMO-2. It is seen that for almost all dihedral angles these three contributions are canceled out. In particular for around $\phi = 90^\circ$, contributions from HOMO and HOMO-1 are canceled. In other regions, the contribution from the HOMO is much smaller than the other contributions and cancellation occurs for contributions from HOMO-1 and HOMO-2. For any $\phi$, the contribution from the HOMO-1 (HOMO-2) is very large. The contribution from the HOMO is only large around $\phi = 90^\circ$ in this region, this contribution is much larger than the total contribution. Any specific relation between the spectrum of $E_{PV}$ in 2a and 2b states and the contribution from a specific orbital, while correspondence between the contribution from the HOMO (and HOMO-1) and the $E_{PV}$ spectrum in 1a and 1b states is seen clearly.

In our previous work [8], it was proposed that the enhancement of $E_{PV}$ by electronic excitation can be roughly estimated by the $E_{PV}$ contribution from the orbital from which the electron is excited. In the present work, the effectiveness of this estimate is clarified. The $E_{PV}$ spectrum in 1a and 1b states can be understood in this estimate. In the 1a and 1b excited states, the electron in the HOMO is excited to unoccupied orbitals. Hence, if contributions from unoccupied orbitals in the ground state are small in the 1a and 1b states, the value of $E_{PV}$ in the 1a and 1b states can roughly be estimated with the assumption that the contribution from the HOMO ($M_{PV,HOMO}^X$) is simply lost. Actually, this rough estimate explains the value of the 1a and 1b states at $\phi = 90^\circ$ [8]. (The HOMO contribution in Table XI in Ref. [8] is the contribution from two electrons in the HOMO, and half of the HOMO contribution is compared to $E_{PV}$ in excited states.) $E_{PV}$ decreases as angle $\phi$ goes away from $90^\circ$, which corresponds to the decrease of $M_{PV,HOMO}^X$. On the other hand, this estimate does not predict the value of the 2a and 2b states. The enhancement in these excited states is not so simple, since the variation of orbitals...
from HF orbitals by excitation is important as discussed below.

The enhancement in the first excited state is considered to be a special case of enhancement, and cancellation breaking enhancement (CBE) hypothesis is proposed in this paper. If the following four conditions are satisfied, significant PVED enhancement is derived in the first excited state and the enhanced PVED value may be roughly estimated as the $E_{PV}$ contribution from the HOMO,

$$
E_{PV}(\text{CBE}) = - \sum_n \frac{G_F}{2\sqrt{2}} \sqrt{I_{PV,n}} M_{PV,\text{HOMO}}.
$$

Since this can be calculated in only HF or DFT computations, this estimate is considered to be useful. In more accurate forms, it is expressed as $- \sum_n \sum_{i} \frac{G_F}{2\sqrt{2}} \sqrt{I_{PV,n} M_{PV,i}}$, or $G_F \sum_n \sqrt{I_{PV,n}} (\sum_{i} \sqrt{I_{PV,n} M_{PV,i}} - \sum_{i} \sqrt{I_{PV,n} M_{PV,i}})$), where $r_{i}^{O}$ is the excitation ratio of the electron in the i-th occupied orbital and $r_{i}^{U}$ is the excitation ratio to the i-th unoccupied orbital. These improved estimates require a post-SCF calculation to determine the ratios and are much larger than the former simple estimate. The four conditions to be satisfied for deriving significant PVED enhancement are listed below.

1. The $E_{PV}$ contribution from the HOMO is much larger than the total value of $E_{PV}$.
2. In the first excited state, the electron in the ground state HOMO is dominantly excited.
3. Occupied orbitals in the ground state are not significantly modified by the excitation.
4. Contributions from unoccupied orbitals to $E_{PV}$ are small.

The first condition requires that the cancellation between contributions is related to the HOMO. Hence, this condition must be satisfied for enhancement by cancellation breaking. It is not clear whether this condition is satisfied for most chiral molecules. However, many molecules, such as CHFClBr, alanine, serine, and valine, satisfy this condition and this condition can easily be checked by HF or DFT computations. The second and third conditions are satisfied for many molecules, at least for vertical excitation of the molecules studied in this work. If the third condition is not satisfied, post-SCF computations are necessary to determine contributions to $E_{PV}$ from occupied orbitals in excited states. If the modification of occupied orbitals is not small, we can hardly predict the enhancement from the electronic structure in the ground state. In the first excited state, excitation is dominated by the electron in the HOMO and the modification of occupied orbitals is considered to be small. Hence, the second and third conditions are usually satisfied. Similar
of the CBE hypothesis is confirmed for these molecules. 

Small deviations from $E_{PV}$ are seen for heavier elements (Se and Te). The cause of such deviations is discussed later. For other excited states, even the improved estimate cannot correctly predict $E_{PV}$ and the CBE hypothesis is not related to the enhancement in these excited states. In the 2a and 2b states of H$_2$O$_2$ and H$_2$S$_2$, $\gamma$$_{HOMO}$ $\lesssim$ 0.7 for $\phi$ = 60 – 120 $^\circ$, and the second condition of the CBE hypothesis is not satisfied. (To check the CBE hypothesis, the second condition was studied also for the second excited states.) In the 2a and 2b states of H$_2$Se$_2$ and H$_2$Te$_2$, $r_{PV,HOMO}^O$ $\sim$ 0.9 except for $\phi$ $\sim$ 90 $^\circ$, where $r_{PV,HOMO}^O$ $\sim$ 0.2 – 0.4, and the second condition of the CBE hypothesis is satisfied except for $\phi$ $\sim$ 90 $^\circ$.

To confirm that the effect of unoccupied orbitals is small, $f_r(X)$ $\sum_i$ (2$M_{PV,i}$) – $\sum_i$ (2$M_{PV,i}$)) is shown in Fig. 4. In this estimate, contributions from virtual orbitals (to which electrons are excited) are added to the improved estimate in Fig. 3. The difference between Figs. 3 and 4 is negligibly small for the 1a and 1b excited state of all H$_2$X$_2$. Hence, in these molecules, the effect of unoccupied orbitals on the PVED is confirmed to be negligible. This is just the condition required in the CBE hypothesis. There is hope that the effect of unoccupied orbitals is small in other molecules as well. Even in second excited states, a difference between Figs. 3 and 4 is not seen. Hence, the reason why the CBE hypothesis cannot be applied to the second excited states is specu-
FIG. 3. Estimate of $E_{PV}$ in the CBE hypothesis as well as the improved estimate (IE), 
$$-\sum_{i}^{\text{occ}} G_{\text{X}} \sqrt{f_{i}^{\text{X}}} r_{i}^{\text{O}} (2M_{\text{PV},i}) - \sum_{i}^{\text{occ}} r_{i}^{\text{O}} (2M_{\text{PV},i}).$$

FIG. 4. Improved estimate of $E_{PV}$ with contributions from unoccupied orbitals in the ground state, 
$$\sum_{i}^{\text{occ}} G_{\text{X}} \sqrt{f_{i}^{\text{X}}} r_{i}^{\text{O}} (2M_{\text{PV},i}) - \sum_{i}^{\text{occ}} r_{i}^{\text{O}} (2M_{\text{PV},i}).$$
lated to be because the modification of orbitals from HF orbitals in the ground state is not negligible. That is, the third condition is not satisfied. In $H_2X_2$, the fourth condition is negligible, while for other molecules, it is not known whether this condition is satisfied. In the next section, CHFCIBr-type molecules are taken as test molecules and the CBE hypothesis is checked. Before this check, we digress from the CBE hypothesis, and the electron chirality distribution of each orbital is studied for $H_2X_2$.

The electron chirality density near a Te nucleus in the ground state of $H_2Te_2$ is shown in Fig. 5. This result is derived by HF computations. The density is shown in the $yz$ plane, where the internuclear axis is the $y$ axis. All atoms are on the $yz$ plane at $\phi = 180^\circ$, while at $\phi = 0^\circ$, all atoms are on the $xy$ plane. Our result is consistent with the result in Ref. [31]. At $\phi = 0^\circ$, the molecular structure is achiral, the electron chirality density is zero at the position of the Te nucleus. The electron chirality has a $p$-orbital-like distribution and positive and negative regions are rotated by $90^\circ$. The common node point for both regions is located on the Te nucleus. For larger $\phi$, the region with positive chirality is extended and the electron chirality on the Te nucleus becomes nonzero. Around $\phi = 90^\circ$, the node of this positive chirality region appears on the nucleus, and $\lambda_{PV}^{Te} \sim 0$.

Figure 6 shows the contribution from the HOMO to the electron chirality density near the Te nucleus in the electronic ground state of $H_2Te_2$ by HF computations. For $\phi = 0 - 75^\circ$, the electron chirality density is almost zero at the nucleus position, which is on the boundary between negative and positive regions. In comparison between $\phi = 75^\circ$ and $105^\circ$, the negative and positive regions are reversed. Around $\phi = 90^\circ$, the transition occurs by the downward transfer of the positive region. In the contribution from the HOMO-1 (Fig. 7), the region with negative chirality moves downward. Hence, it is seen that a small $E_{PV}$ is realized by the cancellation between contributions from HOMO and HOMO-1. It is speculated that in electronic excited states (1a and 1b), this contribution from the HOMO is removed and the cancellation breaking produces large $E_{PV}$.

### B. CHFCIBr, CHFCII, and CHFBrI

To check the CBE hypothesis for other molecules, the CHFCIBr molecule is studied in this section. This molecule was actually used in a PVED observational experiment [2]. In addition to CHFCIBr, CHFCII and CHFBrI are also studied. These two molecules are derived by replacing the Cl or Br atom with an I atom.

Before calculating $E_{PV}$ in excited states, it should be checked whether these molecules satisfy the conditions of the CBE hypothesis. First, the contribution to $E_{PV}$ from the HOMO is confirmed larger by far than the total $E_{PV}$. In Fig. 5, the contribution to $E_{PV}$ from each Kramers pair in the ground state and accumulated value are shown. These contributions are computed by the HF method. In CHFCIBr, the accumulated value of $E_{PV}$ in CHFCIBr is $-6.82 \times 10^{-18}$ Hartree, the contribution from one electron of the HOMO is $-7.71 \times 10^{-17}$ Hartree. The contribution from the HOMO is about 11 times as large as the total $E_{PV}$. The first condition of the CBE hypothesis is satisfied for CHFCIBr. For CHFCII and CHFBri, the accumulated value of $E_{PV}$ is $-6.64 \times 10^{-17}$ and $-1.56 \times 10^{-16}$ Hartree, respectively, while the contribution from the HOMO is $-1.74 \times 10^{-16}$ and $-5.78 \times 10^{-16}$ Hartree, respectively. The contribution from the HOMO to $E_{PV}$ is much larger than the accumulated $E_{PV}$ for both molecules, and the ratio of the HOMO contribution to the total $E_{PV}$ is about 3 (CHFCII) and 4 (CHFBrI). These two ratios are smaller than that of CHFCIBr, and it is speculated that the enhancement of $E_{PV}$ in CHFCII and CHFBrI is smaller than that in CHFCIBr. Contributions from other valence electrons are much larger than the contribution from the HOMO and even small changes in these orbitals may affect the total $E_{PV}$.

The result of CHFCIBr in the ground state is consistent with values based on HF computations in previous works [32, 33]. Contributions to $E_{PV}$ from Kramers pairs was also reported in Ref. [32]. The trend in the previous work is the same as observed in our result, while the values of HOMO and HOMO-1 are significantly smaller than our values. Their value is about 1/3 compared to our result. Nevertheless, even their small HOMO value is much larger than the total $E_{PV}$.

The second and third conditions are considered to be satisfied and, for the first excited state, we confirmed that electron excitation dominantly comes from the HOMO in our EOM-CCSD computations. Hence, the fourth condition that the matrix element of $\sum_n g_n^C M_{PV,i}^n$ is small for unoccupied orbitals, is checked. For the first and second excited states, contributions to $E_{PV}$ from unoccupied orbitals are much smaller than that from the HOMO. Most contributions are less than 1/10 of the HOMO contribution. Only the lowest unoccupied molecular orbital (LUMO) contributions in the first and second excited states of CHFCII are larger than 1/10 of the HOMO contribution and about 1/6 and 1/7 of the HOMO contribution, respectively. Therefore, the conditions of the CBE hypothesis are satisfied.

Since the conditions of the CBE hypothesis are satisfied, the PVED enhancement is studied for these molecules. The PVED in excited states calculated by the EOM-CCSD method is shown in Tables I, II and III. The adopted values of $\lambda^n$ are $\lambda^C = 10^{-1}$ (a.u.), $\lambda^l = 10^{-3}$ (a.u.), $\lambda^{Cl} = 10^{-3}$ (a.u.), $\lambda^{Br} = 10^{-3}$ (a.u.), and $\lambda^I = 10^{-5}$ (a.u.). In the result of CHFBrI, the contribution of the F atom was not included, since the EOM-CCSD computation with the F atom perturbation did not converge. Since contributions from the F atom are subdominant in Z-vector calculations of both CHFCIBr and CHFCII, the neglected contribution from the F atom can safely be dropped in CHFBrI. For all molecules, the CBE enhancement of $E_{PV}$ is observed in the first ex-
FIG. 5. Distribution of electron chirality density near a Te nucleus in the ground state of H$_2$Te$_2$.

FIG. 6. Distribution of the HOMO contribution to the electron chirality density near a Te nucleus in the ground state of H$_2$Te$_2$. 
TABLE I. $E_{PV}$ in the CHFCIBr molecule. In the rightmost column, the CBE hypothesis prediction is shown and this computation is based on the contribution from the HOMO in the HF computation.

| Atom | $E_{PV}/E_h$ (Z-vector) | $E_{PV}/E_h$ (FFPT) | $E_{PV}/E_h$ (HF) | CBE |
|------|-------------------------|---------------------|-------------------|-----|
| Br   | $-6.78 \times 10^{-18}$ | $-6.85 \times 10^{-18}$ | $1.10 \times 10^{-16}$ | $-1.28 \times 10^{-16}$ | $3.94 \times 10^{-17}$ |
| Cl   | $2.46 \times 10^{-18}$  | $2.48 \times 10^{-18}$  | $1.10 \times 10^{-16}$ | $1.18 \times 10^{-16}$ | $4.00 \times 10^{-17}$ |
| F    | $-7.00 \times 10^{-19}$ | $-6.96 \times 10^{-19}$ | $-5.47 \times 10^{-18}$ | $-5.47 \times 10^{-18}$ | $-2.62 \times 10^{-18}$ |
| C    | $-3.80 \times 10^{-20}$ | $-3.80 \times 10^{-20}$ | $1.14 \times 10^{-19}$ | $-8.14 \times 10^{-19}$ | $4.18 \times 10^{-19}$ |
| Total| $-5.05 \times 10^{-18}$ | $-5.10 \times 10^{-18}$ | $2.15 \times 10^{-16}$ | $-1.58 \times 10^{-17}$ | $7.71 \times 10^{-17}$ |

TABLE II. $E_{PV}$ in the CHFCII molecule. In the rightmost column, the CBE hypothesis prediction is shown and this computation is based on the contribution from the HOMO in the HF computation.

| Atom | $E_{PV}/E_h$ (Z-vector) | $E_{PV}/E_h$ (FFPT) | $E_{PV}/E_h$ (HF) | CBE |
|------|-------------------------|---------------------|-------------------|-----|
| I    | $-4.96 \times 10^{-17}$ | $-4.80 \times 10^{-17}$ | $8.44 \times 10^{-16}$ | $-8.71 \times 10^{-16}$ | $1.37 \times 10^{-16}$ |
| Cl   | $7.36 \times 10^{-18}$  | $7.32 \times 10^{-18}$  | $7.70 \times 10^{-17}$ | $7.86 \times 10^{-17}$ | $3.84 \times 10^{-17}$ |
| F    | $-2.15 \times 10^{-18}$ | $-2.12 \times 10^{-18}$ | $-4.54 \times 10^{-18}$ | $-4.46 \times 10^{-18}$ | $-2.29 \times 10^{-18}$ |
| C    | $-1.25 \times 10^{-19}$ | $-1.25 \times 10^{-19}$ | $4.53 \times 10^{-19}$ | $-2.40 \times 10^{-19}$ | $3.50 \times 10^{-19}$ |
| Total| $-4.45 \times 10^{-17}$ | $-4.32 \times 10^{-17}$ | $9.16 \times 10^{-16}$ | $-8.02 \times 10^{-16}$ | $1.74 \times 10^{-16}$ |

TABLE III. $E_{PV}$ in the CHFBrI molecule. In the rightmost column, the CBE hypothesis prediction is shown and this computation is based on the contribution from the HOMO in the HF computation.

| Atom | $E_{PV}/E_h$ (Z-vector) | $E_{PV}/E_h$ (FFPT) | $E_{PV}/E_h$ (HF) | CBE |
|------|-------------------------|---------------------|-------------------|-----|
| I    | $-1.91 \times 10^{-16}$ | $-1.84 \times 10^{-16}$ | $-2.04 \times 10^{-16}$ | $1.19 \times 10^{-16}$ | $2.12 \times 10^{-16}$ |
| Br   | $9.56 \times 10^{-17}$  | $9.54 \times 10^{-17}$  | $1.06 \times 10^{-15}$ | $1.15 \times 10^{-15}$ | $3.67 \times 10^{-16}$ |
| F    | $-1.50 \times 10^{-18}$ | —                   | —                  | —    | $-1.21 \times 10^{-18}$ |
| C    | $-9.42 \times 10^{-20}$ | $-9.45 \times 10^{-20}$ | $-2.25 \times 10^{-19}$ | $-6.46 \times 10^{-19}$ | $2.18 \times 10^{-19}$ |
| Total| $-9.67 \times 10^{-17}$ | $-8.91 \times 10^{-17}$ | $8.60 \times 10^{-16}$ | $1.26 \times 10^{-15}$ | $5.78 \times 10^{-16}$ |
excited state, and the enhancement ratios are -42 (CHFCIBr), -21 (CHFCII), and -9 (CHFBrI). The observed enhancement of these molecules is weaker than HClBr), -21 (CHFClI), and -9 (CHFBrI). The enhancement in the first excited state is much larger than the second excited state, while for CHFClI and CHFBrI the enhancement of two excited states is comparable. The smallness of the second excited state in CHFCIBr is due to the cancellation between the contributions of Br and Cl atoms. The first and second excited states of these three molecules are in the same spin triplet. The excitation energies of the first and second excited states are 4.960 and 4.961 eV (CHFCIBr), 3.667 and 3.667 eV (CHFCII), and 3.577 and 3.578 eV (CHFBrI), respectively.

As noted in Sec. III, a loose convergence threshold is used for EOM-CCSD computations of excited states. Since the energy difference by the perturbation is $\lambda^n M_{PV}$, computations should be sufficiently accurate to include this effect. If this effect is correctly included, the calculated energy difference, $\Delta E^n = E^n(\lambda^n) - E^n(-\lambda^n)$, is independent of $\lambda^n$. To check this, $\Delta E^n$ of CHFCIBr and CHFCII are shown in Fig. 9. For CHFBrI, $E^n$ are calculated in a single $\lambda^n$ value speculated from computations of CHFCIBr and CHFCII to save computational costs. For our chosen values of $\lambda^n$, all $\Delta E^n$ are proportional to $\lambda^n$. This result clearly confirms that our loose con-
TABLE IV. Dependence of $E_{PV}$ on perturbation parameter $\lambda$ within the FFPT method in the ground (a), first excited (1a), and second excited (2a) electronic states of the CHFClBr molecule. Adopted values are emphasized by italic letters.

| Atom | $\lambda$ (a.u.) | $E_{PV}$ | $E_{a,b}$ (FFPT) |
|------|-----------------|----------|------------------|
|      |                 | a        | 1a               | 2a               |
| Br   | $1.0 \times 10^{-3}$ | $-6.85 \times 10^{-18}$ | $1.10 \times 10^{-16}$ | $-1.28 \times 10^{-16}$ |
|      | $1.0 \times 10^{-4}$ | $-6.66 \times 10^{-18}$ | $1.07 \times 10^{-16}$ | $-1.25 \times 10^{-16}$ |
|      | $1.0 \times 10^{-5}$ | $-1.32 \times 10^{-17}$ | $1.01 \times 10^{-16}$ | $-1.32 \times 10^{-16}$ |
| Cl   | $1.0 \times 10^{-2}$ | $2.48 \times 10^{-18}$ | $1.10 \times 10^{-16}$ | $1.18 \times 10^{-16}$ |
|      | $1.0 \times 10^{-3}$ | $2.48 \times 10^{-18}$ | $1.10 \times 10^{-16}$ | $1.18 \times 10^{-16}$ |
|      | $1.0 \times 10^{-4}$ | $2.96 \times 10^{-18}$ | $1.10 \times 10^{-16}$ | $1.18 \times 10^{-16}$ |
| F    | $1.0 \times 10^{-2}$ | $-6.96 \times 10^{-19}$ | $-5.47 \times 10^{-18}$ | $-5.48 \times 10^{-18}$ |
|      | $1.0 \times 10^{-3}$ | $-6.96 \times 10^{-19}$ | $-5.47 \times 10^{-18}$ | $-5.47 \times 10^{-18}$ |
|      | $1.0 \times 10^{-4}$ | $-9.11 \times 10^{-19}$ | $-5.64 \times 10^{-18}$ | $-5.65 \times 10^{-18}$ |
| C    | $1.0 \times 10^{-4}$ | $-3.80 \times 10^{-20}$ | $1.14 \times 10^{-19}$ | $-8.14 \times 10^{-19}$ |
|      | $1.0 \times 10^{-2}$ | $-3.76 \times 10^{-20}$ | $1.04 \times 10^{-19}$ | $-7.98 \times 10^{-19}$ |
|      | $1.0 \times 10^{-3}$ | $-3.67 \times 10^{-20}$ | $1.08 \times 10^{-19}$ | $-7.73 \times 10^{-19}$ |
|      | $1.0 \times 10^{-4}$ | $-2.62 \times 10^{-20}$ | $1.05 \times 10^{-19}$ | $-9.53 \times 10^{-19}$ |

TABLE V. Dependence of $E_{PV}$ on perturbation parameter $\lambda$ within the FFPT method in the ground (a), first excited (1a), and second excited (2a) electronic states of the CHFClII molecule. Adopted values are emphasized by italic letters.

| Atom | $\lambda$ (a.u.) | $E_{PV}$ | $E_{a,b}$ (FFPT) |
|------|-----------------|----------|------------------|
|      |                 | a        | 1a               | 2a               |
| I    | $1.0 \times 10^{-4}$ | $-5.17 \times 10^{-17}$ | $8.88 \times 10^{-16}$ | $-9.27 \times 10^{-16}$ |
|      | $1.0 \times 10^{-5}$ | $-4.80 \times 10^{-17}$ | $8.44 \times 10^{-16}$ | $-8.71 \times 10^{-16}$ |
|      | $1.0 \times 10^{-6}$ | $-1.74 \times 10^{-16}$ | $6.89 \times 10^{-16}$ | $-1.03 \times 10^{-15}$ |
| Cl   | $1.0 \times 10^{-3}$ | $7.32 \times 10^{-18}$ | $7.70 \times 10^{-17}$ | $7.86 \times 10^{-17}$ |
|      | $1.0 \times 10^{-4}$ | $7.62 \times 10^{-18}$ | $7.72 \times 10^{-17}$ | $7.89 \times 10^{-17}$ |
| F    | $1.0 \times 10^{-3}$ | $-2.12 \times 10^{-18}$ | $-4.54 \times 10^{-18}$ | $-4.46 \times 10^{-18}$ |
|      | $1.0 \times 10^{-4}$ | $-2.32 \times 10^{-18}$ | $-4.79 \times 10^{-18}$ | $-4.71 \times 10^{-18}$ |
| C    | $1.0 \times 10^{-4}$ | $-1.25 \times 10^{-19}$ | $4.53 \times 10^{-19}$ | $-2.40 \times 10^{-19}$ |
|      | $1.0 \times 10^{-2}$ | $-1.24 \times 10^{-19}$ | $4.28 \times 10^{-19}$ | $-2.09 \times 10^{-19}$ |
|      | $1.0 \times 10^{-3}$ | $-1.22 \times 10^{-19}$ | $4.22 \times 10^{-19}$ | $-2.08 \times 10^{-19}$ |
|      | $1.0 \times 10^{-4}$ | $-2.62 \times 10^{-19}$ | $2.54 \times 10^{-19}$ | $-3.37 \times 10^{-19}$ |

Convergence computations are sufficiently accurate to derive $E_{PV}$. Tables IV and V show that the calculated $E_{PV}$ is independent of $\lambda^a$ around our adopted value of $\lambda^a$.

Since the conditions of the CBE hypothesis are satisfied and the enhancement of $E_{PV}$ is confirmed, the accuracy of the simple estimate of $E_{PV}$ by the CBE hypothesis should be checked. In Tables III and IV the prediction of $E_{PV}$ based on the CBE hypothesis is also shown. The CBE hypothesis can successfully predict the existence of enhancement and estimate the order of magnitude of $E_{PV}$. However, its accuracy is significantly worse compared to that calculated for $H_2X_2$. The predicted values of $E_{PV}$ are $1/3, 1/5, and 2/3$ of the values derived by EOM-CCSD for CHFClBr, CHFClII, and CHFBrI in the first excited state, respectively. This gap is not narrowed significantly, even with the usage of the improved formulas. Hence, the absence of effects of other HF orbitals in the CBE prediction is not an important factor of this inaccuracy, and the effect of the modification of some occupied orbitals is speculated to be important. The contributions from HOMO-2 and HOMO-3 are much larger than the HOMO contribution as seen in Fig. 8. Even the small change of an orbital with a large contribution may give a large change of $E_{PV}$. In other words, if the contribution of the HOMO is not a dominant part of $E_{PV}$, the accuracy of the CBE prediction (Eq. 6) may be insufficient despite the successful prediction of the existence of enhancement.

Lastly, the dependence on computational conditions is studied using CHFClBr as an example. The value of $E_{PV}$ in the ground state is strongly dependent on the choice of basis sets. In the CCSD computation with aug-cc-pVTZ, cc-pVTZ, and aug-cc-pVDZ basis sets [34], $E_{PV} = -2.54 \times 10^{-18}, -1.83 \times 10^{-18}$, and $-1.60 \times 10^{-18}$.
In this paper, we have studied the $E_{PV}$ enhancement in electronic excited states for H$_2$X$_2$ ($X = O, S, Sc, Te$), CHFCIBr, CHFCII, and CHFBri. The enhancement of $E_{PV}$ of H$_2$X$_2$ in excited states is highly dependent on the dihedral angle. Through the study of the dihedral angle dependence, the CBE hypothesis has been proposed. This hypothesis has been confirmed for excited states of CHFCIBr, CHFCII, and CHFBri. If the contribution from the HOMO to $E_{PV}$ in the ground state is larger than the sum of those from all occupied orbitals, the CBE hypothesis predicts the existence of the PVED enhancement in the first excited state. Moreover, if the contribution from the HOMO to $E_{PV}$ dominates over other contributions, the CBE hypothesis can predict the value of $E_{PV}$ in the first excited state.

We believe that PVED enhancement by the CBE hypothesis occurs for many other chiral molecules. This enhancement mechanism may be a hint for choosing a chiral molecule for future experiments used to observe PVED. For example, the measurement of the vibrational frequency difference between two enantiomers of a chiral molecule in excited states may be a candidate for future experiments. The precise measurement of the excitation energy difference between an enantiomeric pair is also confirmation of the existence of PVED. In this paper, molecular structures of CHFCIBr, CHFCII, and CHFBri in excited states are chosen to be the same as the ground state. Nevertheless, these molecules are not considered to be achiral in excited states. In optimized structures, the PVED may not be enhanced significantly, but we expect that large PVED enhancement is realized in the structures. The investigation of the PVED in the optimized structure of an excited state is important for actual experimental planning. For a molecule whose structure in the first excited state is not so different from that in the ground state, the PVED enhancement in the CBE hypothesis is available for future experiments.

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