

An *ab initio* study on the ground and low-lying doublet electronic states of SbO₂

Edmond P. F. Lee^{a),b)}

Department of Applied Biology and Chemical Technology, The Hong Kong Polytechnic University, Hung Hom, Hong Kong

John M. Dyke

School of Chemistry, University of Southampton, Highfield, Southampton SO17 1BJ, United Kingdom

Foo-Tim Chau^{a),c)}

Department of Applied Biology and Chemical Technology, The Hong Kong Polytechnic University, Hung Hom, Hong Kong

Wan-Ki Chow

Department of Building Services Engineering, The Hong Kong Polytechnic University, Hung Hom, Hong Kong

Daniel K. W. Mok

Department of Applied Biology and Chemical Technology, The Hong Kong Polytechnic University, Hung Hom, Hong Kong

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Geometry optimization and harmonic vibrational frequency calculations have been carried out on the low-lying doublet electronic states of antimony dioxide (SbO₂) employing a variety of *ab initio* methods, including the complete active space self-consistent field/multireference configuration interaction and the RCCSD(T) methods. Both large and small core relativistic effective core potentials were used for Sb in these calculations, together with valence basis sets of up to aug-cc-pV5Z quality. Contributions from outer core correlation and off-diagonal spin-orbit interaction to relative electronic energies have been calculated. The ground electronic state of SbO₂ is determined to be the \tilde{X}^2A_1 state, as is the case for dioxides of other lighter group 15 *p*-block (or group VA) elements. However, the \tilde{A}^2B_2 and \tilde{B}^2A_2 states are estimated to be only 4.1 and 10.7 kcal/mole above the \tilde{X}^2A_1 state, respectively, at the complete basis set limit. Reliable vertical excitation energies from the \tilde{X}^2A_1 state to low-lying excited states of SbO₂ have been computed with a view to assist future spectral assignments of the absorption and/or laser-induced fluorescence spectra of SbO₂, when they become available. © 2006 American Institute of Physics.

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INTRODUCTION

In recent work, clusters of antimony and antimony oxide have been studied in the gas phase using the technique of molecular beam mass spectrometry.¹⁻⁵ In addition to investigating the physical and chemical properties of matter as it changes from atomic or molecular to the bulk via cluster formation, these studies also attempt to understand the complex mechanism of catalytic oxygen transfer on an antimony surface by using as a model the formation of antimony oxide clusters from the reaction between antimony metal clusters and molecular oxygen in the gas phase. The importance of antimony oxide in catalysis is well known (see, for example, Refs. 3 and 6-8), and there are various industrial applications of antimony oxide relating to its catalytic properties (see, for example, Refs. 3 and 9, and references therein),

such as its use in antimony-halogen synergistic flame retardants in polymeric¹⁰⁻¹² and nanocomposite^{13,14} materials. Unfortunately, the experimental technique of molecular beam mass spectrometry does not provide structural information (geometrical, molecular, and/or electronic) on the detected clusters. In this connection, density functional theory (DFT) and/or *ab initio* calculations carried out on antimony oxide clusters would be invaluable in assisting the interpretation of, and the extraction of the maximum information from, the mass spectrometric results obtained. In fact, such calculations have been reported in some of the above-mentioned mass spectrometric studies.^{3,5} In addition, DFT and/or *ab initio* calculations have been performed in other spectroscopic studies of various antimony oxides in order to assist spectral assignments,^{9,15} and there are also numerous independent computational studies on antimony oxides,¹⁶⁻²² though most of these computational studies are on the diatomic antimony monoxide SbO.¹⁶⁻²¹ To our knowledge, however, there are only two computational studies, which have considered the triatomic antimony dioxide (SbO₂).

^{a)}Authors to whom correspondence should be addressed.

^{b)}Also at School of Chemistry, University of Southampton. Electronic mail: epl@soton.ac.uk

^{c)}Electronic mail: bcfchau@polyu.edu.hk

TABLE I. Basis sets used for Sb.

Basis	ECP ^a	Valence set ^a	Augmented ^b	Frozen ^c	Correlated ^d	Nb ^e
A	ECP28MDF	Aug-cc-pVTZ		4s4p4d	5s ² 5p ³	147
B	ECP46MWB	Aug-cc-pVQZ			5s ² 5p ³	239
B1	ECP46MWB	Aug-cc-pVQZ(<i>spd</i>) ^f			5s ² 5p ³	185
C	ECP28MDF	Aug-cc-pV5Z		4s4p4d	5s ² 5p ³	393
D	ECP28MDF	Aug-cc-pV5Z	3d2f2g2h	4s4p	4d ¹⁰ 5s ² 5p ³	462
E	ECP28MDF	Aug-cc-pV5Z	3s3p3d3f2g2h		4s ² 4p ⁶ 4d ¹⁰ 5s ² 5p ³	481

^aStandard effective core potentials and valence basis sets for Sb are published: the ECP46MWB_aug-cc-pVQZ and ECP28MDF_aug-cc-pV5Z basis sets are from MOLPRO basis set library. The ECP46MWB_aug-cc-pVQZ basis set is the same as the SDB-aug-cc-pVQZ basis from Ref. 27. The ECP28MDF_aug-cc-pVXZ ($X=T$ and 5) basis sets are the same as aug-cc-pVXZ-PP ($X=T$ and 5) basis sets, respectively, from the EMSL basis set library (Ref. 29). For the ECP46MWB and ECP28MDF ECPs, see Refs. 31 and 28. The corresponding aug-cc-pVXZ ($X=T$, Q, and 5) basis sets were used for O and the O $1s^2$ electrons are frozen in all the correlation calculations.

^bAugmented basis functions designed in this work.

^cAlthough these shells of Sb are accounted for by the basis sets used, they are frozen in the correlation calculations. The O $1s^2$ electrons are frozen in all correlation calculations.

^dThese Sb electrons are correlated. The O $2s^22p^4$ electrons are correlated in all correlation calculations.

^eTotal numbers of contracted Gaussian functions in the basis sets for SbO₂.

^fOnly *s*, *p*, *d*, and *f* functions (i.e., no *g* functions).

Both are density functional studies,^{16,22} though one of them²² has evaluated the atomization energy of SbO₂ at the Hartree-Fock (HF) and second-order Moller-Plesset (MP2) levels employing an optimized bent structure of SbO₂ [$r_e(\text{SbO}) = 1.8849 \text{ \AA}$ and $\theta_e(\text{OSbO}) = 120.34^\circ$] obtained at the BP86/SV(P) level of calculation. In the other density functional study of Ref. 16 on SbO₂, the local-density approximation (LDA) functional was used and the ground electronic state was computed to have a linear structure [$r_e(\text{SbO}) = 1.87 \text{ \AA}$ and $\theta_e(\text{OSbO}) = 180.0^\circ$], with a bent first excited electronic state [$r_e(\text{SbO}) = 2.01 \text{ \AA}$, $r_e(\text{OO}) = 1.51 \text{ \AA}$, and $\theta_e(\text{OSbO}) = 44.1^\circ$] calculated to be 0.48 eV (11.1 kcal/mole) higher in energy. In both Refs. 16 and 22, however, the symmetries of the electronic states obtained have not been reported. For Ref. 22, where only one geometrical structure was reported, it can only be assumed that the reported structure is that of the ground electronic state of SbO₂. In any case, it appears that from these two DFT studies,^{16,22} the ground electronic state of SbO₂ and its geometrical structure are not well established. In fact, these two DFT studies give contradictory results concerning the geometry of the ground electronic state of SbO₂: a bent structure from Ref. 22 and a linear structure from Ref. 16.

As the first step in a project to investigate the geometrical and electronic structures of small antimony oxides and their clusters, we have carried out DFT and high-level *ab initio* calculations on the low-lying electronic states of SbO₂ with the aim of determining unambiguously its ground electronic state and geometrical structure. In addition, reliable vertical excitation energies from the ground electronic state of SbO₂ have been computed with the aim of assisting future spectral identification of SbO₂ in absorption and/or laser induced fluorescence (LIF) spectra yet to be recorded on SbO₂ in the gas phase.

It should be noted that the ground electronic states of the dioxides of other lighter members of the group 15 *p*-block (or group VA) elements, NO₂,²³ PO₂ (see, Ref. 24, and references therein), and AsO₂ (see Ref. 25, and references

therein), are reasonably well established. They all have \tilde{X}^2A_1 states, with a bent C_{2v} structure. In this connection, the linear structure obtained for the ground electronic state of SbO₂ from LDA calculations reported in Ref. 16 is clearly at odds with those of the lighter group 15 analog, and hence its reliability is doubtful. However, Ref. 16 is the only available computational study, which has reported the structures of both the ground state and an excited electronic state of SbO₂. Clearly further computational investigation is required to determine the ground state of SbO₂.

Lastly, to our knowledge, the only spectroscopic study published on excited electronic states of SbO₂ is the chemiluminescence spectrum observed from the reaction of a molecular beam of Sb with the scattering gas O₃.²⁶ A broad and structureless chemiluminescence band observed in the 650–900 nm (ca. 44–32 kcal/mole) region has been assigned to emission from electronically excited SbO₂^{*}, in addition to a fluorescence band observed in the 400–530 nm region, assigned to emission from SbO^{*}.

COMPUTATIONAL DETAILS AND THEORETICAL CONSIDERATIONS

The basis sets employed in the present study for Sb are summarized in Table I. Basis sets A, B, B1, and C are published standard basis sets.^{27,28} Basis set A was obtained from the EMSL basis set library.²⁹ Basis sets B, B1, and C are from the basis set library of the MOLPRO suite of programs.³⁰ It should be noted that basis set B (B1) is the same as the SDB-aug-cc-pVQZ basis set of Ref. 27 (see Table I), while basis sets A and C are the same as the aug-cc-pVTZ-PP and aug-cc-pV5Z-PP basis sets,²⁸ respectively, from the EMSL website.²⁹ The effective core potential (ECP), ECP46MWB, of basis set B (B1) is a quasirelativistic large core ECP, which accounts for the $1s^22s^22p^63s^23p^63d^{10}4s^24p^64d^{10}$ core electrons of Sb (see Ref. 31 for the ECPs used in the present study). The ECP28MDF ECP of basis sets A, C, D, and E is a fully relativistic small core ECP, with the $4s^24p^64d^{10}$ elec-

trons of Sb considered together with the $5s^25p^3$ electrons as valence. However, the standard valence basis sets A and C of Sb, which employ the ECP28MDF ECP described above, only have polarized-valence- X -zeta ($X=T$ or 5) quality functions accounting for the $5s^25p^3$ electrons of Sb, with a single contracted function describing each of the $4s$, $4p$, and $4d$ outer core subshells of Sb. In order to account for the $4s^24p^6$ and $4d^{10}$ outer core electrons of Sb adequately in correlation calculations, basis sets D and E were designed in the present study based on basis set C. Tight uncontracted $3d$ (exponents: 5.6, 2.8, 1.4), $2f$ (2.4, 1.3), $2g$ (2.9, 1.3), and $2h$ (2.9, 1.3) functions, and $3s$ (8.0, 4.0, 2.0), $3p$ (6.8, 3.4, 1.7), $3d$ (6.0, 3.0, 1.5), $3f$ (5.6, 3.0, 1.4) $2g$ (2.9, 1.3), and $2h$ (2.9, 1.3) functions were augmented to basis set C to give basis sets D and E in order to account for the Sb outer core $4d^{10}$ and $4s^24p^64d^{10}$ electrons, respectively. The frozen core and correlated electrons of Sb associated with different basis sets used in the correlation calculations carried out in the present study are summarized in Table I, together with the total number of contracted basis functions (Nb) for SbO₂. [The corresponding aug-cc-pVXZ ($X=T$, Q, and 5) basis sets of O are used and the O $1s^2$ electrons are frozen in all correlation calculations.]

First, MP2 calculations and DFT calculations employing the B3LYP, PW91PW91, and HTCH407 functionals were carried out using basis set A. Geometry optimization and harmonic vibrational frequency calculations were performed using the GAUSSIAN 03 suite of programs.³² The geometries of the linear and bent structures of SbO₂ reported in Refs. 16 and 22 were used as initial guesses. These preliminary calculations are for the purpose of identifying the symmetries of the linear and bent electronic states reported previously.^{16,22}

Based on the results obtained from the MP2 and DFT calculations, it was felt that a more thorough survey on the low-lying electronic states of SbO₂ would be required. At the same time, it was decided to move on from the MP2 and DFT methods, which are based on unrestricted-spin wave functions (using GAUSSIAN 03), to methods which are based on restricted-spin wave functions (using MOLPRO), so as to avoid the problem of spin contamination associated with unrestricted-spin wave functions. Survey complete active space self-consistent field (CASSCF) geometry optimization calculations using basis set B1 were first carried out, followed by RCCSD(T) and/or CASSCF/MRCI (multireference configuration interaction) geometry optimization calculations using the larger basis sets B, C, and/or D. (A full valence active space was employed in all the CASSCF and MRCI calculations; see also footnotes c and d of Table I.) Numerical harmonic vibrational frequency calculations were also carried out on the lowest energy, bent, and linear electronic states of SbO₂ at the RCCSD(T) level using basis set B. Finally, RCCSD(T) single geometry energy calculations using basis set E were also performed at RCCSD(T)/D geometries for some low-lying states of SbO₂, and the geometry of the ground state of SbO₂ was optimized at the RCCSD(T)/E level of calculation. The purpose of this series of calculations is to obtain reliable adiabatic relative electronic energies of the low-lying electronic states of SbO₂ and hence to determine the ground state of SbO₂. In this connection, geometry

optimization calculations at higher levels of theory were only carried out for a few lowest-lying electronic states.

Further calculations were carried out to obtain reliable vertical excitation energies from the ground electronic state of SbO₂. The purpose of these calculations is to assist future spectral assignments of absorption and/or LIF spectra yet to be observed. In relation to assisting spectral assignments, vertical excitation energies are more relevant than adiabatic excitation energies, as the adiabatic position of a spectral band may not be identifiable because of having a small Franck-Condon factor. CASSCF, CASSCF/MRCI, and/or RCCSD(T) calculations using basis sets B1, B; C, and/or D were carried out at appropriate optimized geometries of the ground electronic state of SbO₂ in order to obtain reliable vertical excitation energies.

In the above calculations, scalar relativistic effects from Sb are accounted for by using the quasirelativistic (ECP46MWB) or relativistic (ECP28MDF) ECPs (see, for example, Ref. 27). The use of different valence basis sets and/or different frozen cores for Sb is to assess the effects of basis size extension to the complete basis set (CBS) limit and/or outer core electron correlation from Sb, respectively, on the optimized geometrical parameters and computed relative electronic energies. The computed CASSCF and/or MRCI wave functions, and/or the computed T_1 diagnostics from the RCCSD(T) calculations would provide us information on the importance of multireference character for the low-lying electronic states of SbO₂.

What remains to be considered is spin-orbit (SO) coupling among the low-lying electronic states of SbO₂. Since the ground electronic state of SbO₂ and the two lowest-lying excited states, which are close in energy to the ground state, as determined in the present study, have C_{2v} structures (see next section), and are therefore nondegenerate states, they do not have diagonal spin-orbit splitting. Nevertheless, off-diagonal spin-orbit interactions between states, which are close to each other in energy, may be significant for a molecule containing the heavy fourth row element Sb. Consequently, spin-orbit coupling among low-lying states of SbO₂ was calculated explicitly by the state-interacting method, which employs the average-state CASSCF wave functions of the electronic states considered, as implemented in MOLPRO. Spin-orbit matrix elements and eigenstates were computed using either the Breit-Pauli (BP) operator or spin-orbit pseudopotentials (ECPs). In the former case of all-electron calculations, uncontracted s , p , and d functions from the DZVP-DFT-orb basis sets³³ (which are all-electron basis sets) were used for Sb and O. In the latter case with the ECP28MWB ECP for Sb, uncontracted s , p , and d functions of basis set B were used. In both cases, more reliable computed electronic energies, which include dynamic electron correlation, were used for the diagonal spin-orbit matrix elements, instead of the CASSCF energies associated with the average-state CASSCF wave functions used in the spin-orbit calculations. For vertical excitation energies, the CASSCF/MRCI+D (internally contracted MRCI plus Davidson correction) energies of the states considered, obtained using basis set B, were used instead of the average-state CASSCF energies for the diagonal spin-orbit matrix

elements. Four states (the lowest state of each symmetry in the C_{2v} symmetry group) and eight states (the lowest two states of each symmetry) were considered separately in these calculations. In addition, the RCCSD(T)/D energies of the lowest four states were also used for the diagonal spin-orbit matrix elements in the CASSCF spin-orbit calculations using the ECP basis set. For the computed T_e 's of the first two excited electronic states of SbO_2 , spin-orbit coupling was included at each RCCSD(T)/D minimum-energy geometry, using the ECP basis set mentioned above. The electronic energies of the states considered in these spin-orbit calculations were computed at the RCCSD(T)/B level at the relevant geometry and were used for the diagonal spin-orbit matrix elements. Only the lowest three states of SbO_2 have been considered in these CASSCF SO calculations, as higher states are significantly higher in energy at the geometries concerned.

RESULTS AND DISCUSSION

MP2 and DFT results

First, from Table II, it can be seen that MP2 and DFT calculations give the consistent result that the lowest energy state is the 2A_1 state with the electronic configuration of $\cdots(5a_1)^1(1b_1)^2(4b_2)^2(1a_2)^2$. However, the calculated bond angle of 137.33° obtained at the MP2 level is significantly larger than those (ca. 122°) obtained from DFT calculations with the three functionals used. Second, the electronic state obtained for the linear structure is also consistent from both MP2 and DFT calculations to be a ${}^2\Sigma_g^+$ state. This linear state is calculated to be higher in energy than the lowest 2A_1 state by ca. 13–21 kcal/mole, with the computed DFT relative energies (with respect to the lowest 2A_1 state) larger than the MP2 one. Third, with an initial geometry with a bent structure with a small bond angle of 44.1° as obtained from LDA calculations of Ref. 16, MP2 geometry optimization leads to a 2A_2 state, which has a computed PUMP2 (spin projection of unrestricted-spin MP2 energy) energy of 68.0 kcal/mole higher than the lowest 2A_1 state. However, DFT geometry optimization calculations with this initial geometry with the three functionals used give a 2B_1 state in all cases, and this 2B_1 state has computed energies ranging from 24.7 to 27.9 kcal/mole above the lowest 2A_1 state, depending on the functional used. Comparing these results with those reported previously, it seems clear that the bent structure of Ref. 22 corresponds to the lowest energy 2A_1 state obtained here, while the linear and bent structures of Ref. 16 correspond to the ${}^2\Sigma_g^+$ and 2B_1 states obtained here, respectively. In this connection, both structures from LDA calculations of Ref. 16 are those of excited states of SbO_2 .

When the computed MP2 and DFT harmonic vibrational frequencies obtained in the present study are considered (see Table II), the MP2 calculations give one unrealistically large vibrational frequency for both the 2A_1 state [the asymmetric stretching mode of 2174 cm^{-1} (b_2)] and the ${}^2\Sigma_g^+$ state [the asymmetric stretching mode of 2571 cm^{-1} (σ_u)], as compared with the corresponding DFT values (less than 800 cm^{-1} ; see Table II). It is noted that the computed $\langle S^2 \rangle$ values of the unrestricted-spin Hartree-Fock (UHF) wave

functions for the 2A_1 and ${}^2\Sigma_g^+$ states of SbO_2 obtained from the MP2 calculations are 0.853 and 0.817, respectively, suggesting that spin contaminations in the UHF wave functions of these two doublet states from higher-spin quartet states are not negligible. In contrast, the computed $\langle S^2 \rangle$ values from the corresponding DFT calculations are less than 0.77 in all cases. It appears that the unrealistically large computed asymmetric stretching frequencies of these two states of SbO_2 obtained from the MP2 calculations are very likely due to spin contaminations in the UHF wave functions. In addition, the MP2 vibrational frequencies are all real for the ${}^2\Sigma_g^+$ state, but the DFT calculations give imaginary vibrational frequencies for the doubly degenerate bending (π_u) modes with all three functionals used for this state, indicating that this linear structure is a saddle point on the DFT energy surfaces. In view of the above considerations, it is concluded that the MP2 method is probably inadequate for the low-lying electronic states of SbO_2 . We will come back to this point, when results obtained from higher level calculations are considered.

CASSCF and higher level calculations

The results obtained from CASSCF, RCCSD(T), and CASSCF/MRCI geometry optimization calculations are summarized in Tables II and III. First, from Table II, when the results obtained from the CASSCF/B1 and RCCSD(T)/B calculations are compared, they are qualitatively, reasonably consistent with each other. This is generally the case for both the optimized geometrical parameters and computed relative electronic energies, for almost all of the electronic states of SbO_2 under investigation. The only major qualitative difference is with the $(2) {}^2B_1$ state with the $\cdots(4a_1)^2(2b_1)^1(4b_2)^2(1a_2)^2$ configuration: CASSCF/B1 geometry optimization calculations give a bond angle of 158.9° , but RCCSD(T)/B calculations give a linear structure, which corresponds to a ${}^2\Pi_u$ state. Apparently, dynamic electron correlation favors a larger bond angle for this state. This seems to be also the case for the lowest energy 2A_1 state, though to a lesser extent: CASSCF calculations give a bond angle of 117.1° , while RCCSD(T) calculations give a bond angle of 122.8° . Regarding computed relative electronic energies of the low-lying electronic states of SbO_2 investigated, those obtained from CASSCF calculations, where dynamic electron correlation is lacking, have a narrower energy spread than those from RCCSD(T) calculations. It is clear that dynamic electron correlation is important in determining the relative electronic energy separations among low-lying electronic states of SbO_2 .

Second, when the RCCSD(T)/B results are compared with the MP2 and DFT results, agreement for both optimized geometrical parameters and computed relative electronic energies is generally reasonably good. Nevertheless, one exception should be noted. Although the computed RCCSD(T)/B bond angle for the 2A_1 state of 122.8° is in very good agreement with DFT values of ca. 122.6° obtained using the PW91PW91 and HCTH407 functionals, it is considerably smaller than the MP2 value of 137.3° . This supports the conclusion made above that the MP2 method is

TABLE II. The optimized geometrical parameters (r_e in Å and θ_e in deg), computed harmonic vibrational frequencies (ω_e 's in cm⁻¹), and relative electronic energies (E_{rel} in kcal/mole) of some low-lying electronic states of SbO₂ obtained at different levels of calculation. (For basis sets used, see Table I).

	r_e (SbO)	θ_e	Vibrational frequencies	E_{rel}
Previous studies				
LDA ^a				
Electronic state unknown	2.01 ^b	44.1		11.1 ^b
Electronic state unknown	1.87	180.0		0.0
BP86/SV (P) ^c				
Electronic state unknown	1.8849	120.34		0.0
Present work				
MP2/A				
² A ₂	2.1502 ^d	36.78	457(a ₁), 686(b ₂), 1013(a ₁)	68.0
² Σ _g	1.8243	180.0	225(π _u), 924(σ _g), 2571(σ _u)	13.4
² A ₁	1.8097	137.33	225(a ₁), 904(a ₁), 2174(b ₂)	0.0
B3LYP/A				
² B ₁	1.9986 ^e	44.41	513(a ₁), 568(b ₂), 870(a ₁)	24.7
² Σ _g	1.8658	180.0	294i(π _u), 662(σ _u), 704(σ _g)	20.7
² A ₁	1.8489	120.98	206(a ₁), 578(b ₂), 756(a ₁)	0.0
PW91PW91/A				
² B ₁	2.0140	44.41	496(a ₁), 555(b ₂), 847(a ₁)	26.4
² Σ _g	1.8804 ^f	180.0	270i(π _u), 674(σ _g), 675(σ _u)	17.4
² A ₁	1.8623	122.54	199(a ₁), 687(b ₂), 725(a ₁)	0.0
HCTH407/A				
² B ₁	2.0048	43.90	497(a ₁), 562(b ₂), 849(a ₁)	27.9
² Σ _g	1.8709	180.0	279i(π _u), 670(σ _u), 675(σ _g)	18.0
² A ₁	1.8528 ^g	122.72	202(a ₁), 680(b ₂), 725(a ₁)	0.0
CASSCF/B1				
² B ₁ (4a ₁) ² (2b ₁) ¹ (4b ₂) ² (1a ₂) ²	1.8581	158.91		62.5
² A ₂ (5a ₁) ² (2b ₁) ² (3b ₂) ² (1a ₂) ¹	2.1705 ^h	36.06		49.9
² Σ _g ⁺ (3σ _u) ² (1π _u) ⁴ (1π _g) ⁴ (4σ _g) ¹	1.8802	180.0		24.8
² B ₁ (5a ₁) ² (2b ₁) ¹ (3b ₂) ² (1a ₂) ²	1.9860 ⁱ	46.56		23.3
² A ₂ (5a ₁) ² (1b ₁) ² (4b ₂) ² (1a ₂) ¹	1.8965	100.29		5.2
² B ₂ (5a ₂) ¹ (1b ₁) ² (4b ₂) ¹ (1a ₂) ²	1.8778	89.83		1.7
² A ₁ (5a ₁) ¹ (1b ₁) ² (4b ₂) ² (1a ₂) ²	1.8516	117.06		0.0
RCCSD(T)/B				
² B ₁ (4a ₁) ² (2b ₁) ¹ (4b ₂) ² (1a ₂) ²	1.8430	179.99 ^j		88.0
² A ₂ (5a ₁) ² (2b ₁) ² (3b ₂) ² (1a ₂) ¹	2.1648 ^k	35.76		63.4
² B ₁ (5a ₁) ² (2b ₁) ¹ (3b ₂) ² (1a ₂) ²	1.9758 ^l	45.88		26.7
² Σ _g ⁺ (3σ _u) ² (1π _u) ⁴ (1π _g) ⁴ (4σ _g) ¹	1.8609	180.0	233i(π _u), 691(σ _g), 1434(σ _u)	20.0
² A ₂ (5a ₁) ² (1b ₁) ² (4b ₂) ² (1a ₂) ¹	1.8905	100.96		10.7
² B ₂ (5a ₁) ² (1b ₁) ² (4b ₂) ¹ (1a ₂) ²	1.8686	89.13		5.0
² A ₁ (5a ₁) ¹ (1b ₁) ² (4b ₂) ² (1a ₂) ²	1.8363	122.81	205(a ₁), 762(a ₁), 829(b ₂)	0.0

^aReference 16.

^bThe optimized OO bond length is 1.51 Å (giving a OSbO bond angle of ca. 44.1°) and the relative energy is 0.48 eV.

^cReference 22.

^dThe optimized OO bond length is 1.3570 Å.

^eThe optimized OO bond length is 1.5106 Å.

^fThe optimized OO bond length is 1.5221 Å.

^gThe optimized OO bond length is 1.4990 Å.

^hThe optimized OO bond length is 1.3437 Å.

ⁱThe optimized OO bond length is 1.5698 Å.

^jBecame a linear ²Π_u state.

^kThe optimized OO bond length is 1.3294 Å.

^lThe optimized OO bond length is 1.5402 Å.

TABLE III. The optimized geometrical parameters (r_e in Å and θ_e in deg) and relative electronic energies (E_{rel} in kcal/mole) of some low-lying electronic states of SbO_2 obtained at higher levels of calculation. [See Table I for the basis set and frozen core used and the electrons included in the correlated calculations. Note that with basis set B, the $4s^2 4p^6 4d^{10}$ electrons of Sb are accounted for by the ECP46MWB ECP and hence are not included in the numbering of the occupied molecular orbitals. However, with the basis sets C, D, and E, the $4s^2 4p^6 4d^{10}$ electrons of Sb are included in the Hartree-Fock calculations and hence are included in the numbering of the occupied molecular orbitals. The major configurations, the calculated CI coefficient of the major configuration (c_0) and the sum of the squares of the CI coefficients of the reference configurations (Σc_{ref}^2) obtained from the MRCI calculations, and the computed T_1 diagnostics from the RCCSD(T) calculations are given in the first column for CASSCF/MRCI+D/C calculations, and RCCSD(T)/D and RCCSD(T)/E calculations, respectively.]

RCCSD(T)/B	r_e	θ_e	$E_{\text{rel}}(\text{RCCSD})$	$E_{\text{rel}}\{\text{RCCSD(T)}\}$
${}^2\Sigma_g^+ (3\sigma_u)^2(1\pi_u)^4(1\pi_g)^4(4\sigma_g)^1$	1.8609	180.0	24.60	20.04
${}^2A_2 (5a_1)^2(1b_1)^2(4b_2)^2(1a_2)^1$	1.8905	100.96	8.67	10.69
${}^2B_2 (5a_1)^2(1b_1)^2(4b_2)^1(1a_2)^2$	1.8686	89.13	4.43	5.01
${}^2A_1 (5a_1)^1(1b_1)^2(4b_2)^2(1a_2)^2$	1.8363	122.81	0.0	0.0
RCCSD(T)/C	r_e	θ_e	(RCCSD)	{RCCSD(T)}
${}^2\Sigma_g^+ (1\delta_g)^2(4\sigma_u)^2(2\pi_u)^4(7\sigma_g)^1$	1.87713	180.0	23.01	18.62
${}^2A_2 (9a_1)^2(3b_1)^2(6b_2)^2(2a_2)^1$	1.9077	100.34	8.86	10.62
${}^2B_2 (9a_1)^2(3b_1)^2(6b_2)^1(2a_2)^2$	1.8843	88.09	4.16	4.71
${}^2A_1 (9a_1)^1(3b_1)^2(6b_2)^2(2a_2)^2$	1.8514	122.88	0.0	0.0
CASSCF/MRCI+D/C	r_e	θ_e	(MRCI)	(MRCI+D)
${}^2\Sigma_g^+ c_0=0.869, \Sigma c_{\text{ref}}^2=0.947$	1.8667	180.0	21.64	20.30
${}^2A_2 c_0=0.863, \Sigma c_{\text{ref}}^2=0.949$	1.9004	100.47	8.85	10.02
${}^2B_2 c_0=0.869, \Sigma c_{\text{ref}}^2=0.948$	1.8809	88.95	3.92	4.07
${}^2A_1 c_0=0.879, \Sigma c_{\text{ref}}^2=0.948$	1.8379	120.71	0.0	0.0
RCCSD(T)/D	r_e	θ_e	(RCCSD)	{RCCSD(T)}
${}^2A_2 T_1=0.022$	1.8820	100.31	9.23	10.68
${}^2B_2 T_1=0.033$	1.8597	87.94	3.92	4.50
${}^2A_1 T_1=0.031$	1.8277	121.87	0.0	0.0
RCCSD(T)/E/RCCSD(T)/D	r_e	θ_e	(RCCSD)	{RCCSD(T)}
${}^2A_2 T_1=0.020$			9.20	10.65
${}^2B_2 T_1=0.029$			3.79	4.39
${}^2A_1 T_1=0.027$			0.0	0.0
RCCSD(T)/E	r_e	θ_e	(RCCSD)	{RCCSD(T)}
${}^2A_1 T_1=0.027$	1.8271	121.75		

inadequate for low-lying electronic states of SbO_2 . Higher order electron correlation appears to be important in determining reliably the equilibrium geometrical parameters even for the ground electronic state of SbO_2 . In addition, the computed CASSCF bond angle of 117.1° for the lowest 2A_1 state is much closer to the corresponding DFT and RCCSD(T) values than the MP2 value. This comparison also supports the conclusion made above that spin contamination in the UHF wave function used in the MP2 calculations has an adverse effect on the MP2 results for this state.

Third, harmonic vibrational frequencies have been computed for the lowest 2A_1 state and also the linear ${}^2\Sigma_g^+$ state at the RCCSD(T)/B level of calculation. For the linear ${}^2\Sigma_g^+$ state, imaginary frequencies were obtained for the doubly degenerate bending π_u modes, in agreement with DFT results, but contrary to MP2 results, which have all real vibrational frequencies for this linear state as discussed above. It is concluded that the MP2 energy surface of the ${}^2\Sigma_g^+$ state is unreliable, similar to the conclusion made above. Although the bending region of the electronic energy surface with C_{2v} symmetry near to the linear ${}^2\Sigma_g^+$ state saddle point has not

been investigated in detail, upon bending following the π_u mode with computed RCCSD(T) and/or DFT imaginary frequencies, the ${}^2\Sigma_g^+$ state becomes a 2A_1 state. Since the lowest energy 2A_1 state is calculated to be ca. 20 kcal/mole lower in energy [at the RCCSD(T)/B level of calculation] than the ${}^2\Sigma_g^+$ state saddle point and there is no other 2A_1 state nearby, it is almost certain that the ${}^2\Sigma_g^+$ state saddle point is the barrier of linearity of the lowest 2A_1 state. For the lowest 2A_1 state, the vibrational frequency of the asymmetric stretching b_2 mode computed at the RCCSD(T)/B level is 829 cm^{-1} , which is considerably smaller than the corresponding, rather large and unrealistic, MP2 value of 2174 cm^{-1} , and is definitely closer to the DFT values of $578\text{--}687\text{ cm}^{-1}$ (depending on the functional used) than the MP2 value. However, at the RCCSD(T)/B level, the computed symmetric stretching frequency of 762 cm^{-1} is smaller than the asymmetric stretching frequency of 829 cm^{-1} , but with DFT calculations, the calculated symmetric stretching frequencies (between 725 and 756 cm^{-1}) are larger than the corresponding asymmetric stretching frequencies (between 578 and 678 cm^{-1}) with the three functionals used. Based on results shown in Table II, it

is concluded that DFT results are probably more reliable than MP2 results for the low-lying electronic states of SbO_2 , but, from the theoretical point of view, the RCCSD(T) results should be the most reliable. In this connection, the reliability of the computed harmonic vibrational frequencies obtained from DFT calculations, particularly regarding the order/magnitudes of the symmetric and asymmetric stretching modes, is doubtful (see also the last section).

For calculations performed at higher levels of theory, only a few low energy electronic states of SbO_2 were considered and the results are summarized in Table III. It should be noted that the numbering of the molecular orbitals used in the electronic configurations given in Table III is different with different ECPs used in the calculations. Nevertheless, the electronic configurations and states obtained with the two ECPs (and all basis sets) used are consistent with each other. First, the calculated CI coefficients of the major electronic configuration of the states considered, as obtained from MRCI calculations using basis set C, and the computed T_1 diagnostics obtained from RCCSD(T) calculations using basis sets D and E are given in Table III. Both show that multireference character is insignificant for these states at their equilibrium geometries.

Second, considering the optimized geometrical parameters obtained at different levels of calculation, those calculated using the CASSCF/MRCI+D method are generally smaller than those using the RCCSD(T) method with the same basis set C (see Table III). Since the MRCI method is not size consistent, while the RCCSD(T) method is size consistent, the RCCSD(T) results should be theoretically more reliable, particularly in the present cases, where multireference character for these states of SbO_2 has been shown to be unimportant, as discussed. Therefore, we will focus on the RCCSD(T) results. Using a larger basis set and/or including the $4d^{10}$ and/or $4s^2 4p^6 4d^{10}$ electrons of Sb in the correlation calculation leads to a smaller bond angle for all the bent states studied. In this connection, the computed RCCSD(T)/D [or RCCSD(T)/E, for the lowest energy 2A_1 state] bond angles are probably the most reliable and are upper limits to the true values. The maximum theoretical uncertainty associated with the best computed bond angles is probably around 1° based on the results obtained in the present study. For example, it is essentially the inclusion of the $4d^{10}$ outer core electrons of Sb in the correlation calculation, which reduces the equilibrium bond angle of the $\tilde{X}{}^2A_1$ state of SbO_2 by ca. 1° [the difference between the RCCSD(T)/C and RCCSD(T)/D bond angles; the inclusion of the $4s^2 4p^6$ electrons of Sb with basis set E has insignificantly small effects on the computed equilibrium geometrical parameters]. Regarding the optimized bond lengths, they increase from using basis set B (quadruple-zeta quality) to basis set C (quintuple-zeta quality), but decrease when the $4d^{10}$ electrons of Sb are included in the correlation calculations, using basis set D. In this connection, the best calculated bond lengths are probably close to the corresponding computed RCCSD(T)/B values, with uncertainties of ca. $\pm 0.01 \text{ \AA}$.

Third, considering computed relative electronic energies (T_e) obtained at different levels of calculation, contributions from both the Davidson corrections in the MRCI+D calcu-

lations and the triple excitations in the RCCSD(T) calculations to the computed relative electronic energies are generally small, but not insignificant (mostly less than 2 kcal/mole). The largest contribution from triple excitations in a RCCSD(T) calculation to the computed T_e is with the ${}^2\Sigma_g^+$ state [larger than 4 kcal/mole at the RCCSD(T) level with basis sets B and C; see Table III]. Nevertheless, the computed RCCSD(T) and MRCI+D relative electronic energies obtained with the same basis set C are almost identical, with the only exception being the ${}^2\Sigma_g^+$ state, where the computed MRCI+D/C T_e of 20.30 kcal/mole is larger than the RCCSD(T)/C value of 18.62 kcal/mole by ca. 1.7 kcal/mole. If the ${}^2\Sigma_g^+$ state, which is a saddle point on the RCCSD(T)/B electronic energy surface, is ignored, it can be concluded that the computed RCCSD(T) and CASSCF/MRCI relative electronic energies obtained for the three lowest energy states of SbO_2 (2A_2 , 2B_2 , and 2A_1) with basis sets B and C are very consistent. When outer core correlation is included in the RCCSD(T) calculations, where basis sets D and E were employed, it can be seen that its effects on the computed relative electronic energies of the three lowest states studied are negligibly small (less than 0.5 kcal/mole). Based on the RCCSD(T) results obtained using basis sets B, C, D, and E, the maximum computed uncertainties associated with the computed T_e values of the $\tilde{A}{}^2B_2$ and $\tilde{B}{}^2A_2$ states of SbO_2 obtained at the RCCSD(T)/E/RCCSD(T)/D level should be less than 1 kcal/mole. It should be noted that calculated spin-orbit contributions to the computed electronic energies of the lowest three electronic states of SbO_2 are negligibly small (a decrease of 0.6 cm^{-1} for the $\tilde{X}{}^2A_1$ state, a decrease of 0.8 cm^{-1} for the $\tilde{A}{}^2B_2$ state, and an increase of 0.1 cm^{-1} for the $\tilde{B}{}^2A_2$ state).

Summarizing, the computed relative electronic energies of the three lowest energy states of SbO_2 reported here are found to be very consistent at higher levels of calculation and hence are concluded to be highly reliable. In this connection, the ground electronic state of SbO_2 is determined unambiguously to be the $\tilde{X}{}^2A_1$ state. Nevertheless, the T_e values of the $\tilde{A}{}^2B_2$ and $\tilde{B}{}^2A_2$ states of SbO_2 are estimated to be 4.1 and 10.7 kcal/mole at the complete basis set (CBS) limit [taking half of the difference between the RCCSD(T)/B and RCCSD(T)/E values as the CBS correction]. The $\tilde{A}{}^2B_2$ state of SbO_2 is thus quite close in energy to the $\tilde{X}{}^2A_1$ state.

Vertical excitation energies

The computed vertical excitation energies (T_{vert}) and transition dipole moments (TDMs) obtained from average-state CASSCF calculations using basis set B1 are summarized in Table IV. The results obtained from the four-state and eight-state calculations are shown and their differences, which are small, are primarily due to their different state-average CASSCF wave functions. The computed TDMs suggest that electronic excitation to the $(2) {}^2A_1$ state from the $\tilde{X}{}^2A_1$ state is the strongest, followed by excitation to the $(1) {}^2B_2$ (or $\tilde{A}{}^2B_2$) state. These results on SbO_2 are similar to those reported previously on PO_2 .²⁴ However, the $(2) {}^2A_1$

TABLE IV. Computed vertical excitation energies (T_{vert} in kcal/mole) and transition dipole moment [TDM (the computed transition dipole moment matrix element: (excited state|DMX, Y, or Z [dipole moment in the x , y , or z direction (under TDM in parentheses)]|ground state)) in Debye] of some low-lying doublet states of SbO_2 from the \tilde{X}^2A_1 state (the CASSCF averaged-state calculations of the low-lying excited states of SbO_2 were carried out at the CASSCF/MRCI+D/B optimized geometry of the \tilde{X}^2A_1 state) by four-states [the four states are the lowest state of symmetry belonging to each irreducible representation of the C_{2v} symmetry point group (including the \tilde{X}^2A_1 state)] and eight-states [the eight states are the lowest two states of symmetry belonging to each irreducible representation of the C_{2v} symmetry point group (including the \tilde{X}^2A_1 state)] averaged-state CASSCF calculation employing basis set B1 (see Table I).

CASSCF	States	Major configuration	Four states		Eight states	
			T_{vert}	TDM	T_{vert}	TDM
	2A_2	$(5a_1)^2(1b_1)^2(4b_2)^2(1a_2)^1$	13.7	0.0	13.7	0.0
	2B_2	$(5a_1)^2(1b_1)^2(4b_2)^1(1a_2)^2$	16.5	0.965(Y)	16.3	0.829(Y)
	2B_1	$(5a_1)^2(1b_1)^1(4b_2)^2(1a_2)^2$	50.6	0.747(X)	48.5	0.692(X)
	2A_1	$(4a_1)^1(5a_1)^2(1b_1)^2(4b_2)^2(1a_2)^2$			55.7	1.141(Z)
	2B_2	$(5a_1)^2(1b_1)^2(3b_2)^1(4b_2)^2(1a_2)^2$			59.4	0.455(Y)
	2B_1	$(4a_1)^2(1b_1)^2(2b_1)^1(4b_2)^2(1a_2)^2$			69.9	-0.717(X)
	2A_2	$(5a_1)^1(1b_1)^2(2b_1)^1(4b_2)^1(1a_2)^2$			79.4	0.0

TABLE V. Computed vertical excitation energies (T_{vert} in kcal/mole) of some low-lying doublet states (see Table IV for the main configurations of the electronic states) of SbO_2 from the \tilde{X}^2A_1 state obtained at higher levels of calculation. [Calculations were carried out at the corresponding CASSCF/MRCI+D or RCCSD(T) optimized geometries of the \tilde{X}^2A_1 state of SbO_2 . With basis B, the CASSCF/MRCI calculations have considered two roots for the state symmetries of A_1 , B_1 , B_2 , and A_2 . With basis C, only the lowest 2A_2 state was considered.]

T_{vert}	2A_2	2B_2	2B_1	2A_1	2B_2	2B_1	2A_2
CASSCF/B	17.92	16.98	53.39	57.32	62.72	69.17	78.99
CASSCF/MRCI/B	19.87	20.14	51.94	54.31	60.63	68.32	81.15
CASSCF/MRCI+D/B	19.58	18.96	51.16	53.10	59.72	67.66	81.00
CAS/MRCI+D/B+SO(ecp_4states) ^a	20.16	18.71	51.16				
CAS/MRCI+D/B+SO(ecp_8states) ^b	20.15	18.71	51.16	53.10	59.72	67.66	81.01
CAS/MRCI+D/B+SO(all_e_8states) ^c	20.13	18.70	50.16	53.62	60.23	67.70	81.02
CASSCF/C	13.16	17.26	53.67	57.25	62.70	68.91	
CASSCF/MRCI/C	19.17	18.97	52.36	54.38	60.72	68.01	
CASSCF/MRCI+D/C	20.79	19.1	51.66	53.26	59.89	67.35	
RCCSD ^d /C	19.71	21.10	55.75			66.96	
RCCSD(T)/C	21.99	22.02	52.95			65.81	
RCCSD ^e /D	19.55	20.86	57.29			68.67	
RCCSD(T) ^e /D	21.48	21.61	54.36			66.69	
RCCSD(T)/D+SO(ecp_4states) ^f	21.38	21.71	54.36				

^aThe average-state CASSCF spin-orbit (SO) interaction calculations were carried out at the CASSCF/MRCI+D/B geometry of the \tilde{X}^2A_1 state of SbO_2 , employing the ECP46MWB ECP and the uncontracted s , p , and d functions of basis set B (the total number of basis functions used is 176; see Table I and text). The CASSCF/MRCI+D/B energies were used for the SO diagonal matrix elements. Only the lowest doublet states of the four symmetries were considered. The \tilde{X}^2A_1 state of SbO_2 was lowered by 0.68 cm^{-1} through off-diagonal SO interaction.

^bAs with footnote a, but with eight states, the \tilde{X}^2A_1 state of SbO_2 was lowered by 0.78 cm^{-1} through off-diagonal SO interaction.

^cAs with footnotes a and b, but the basis sets used are the uncontracted s , p , and d functions of the all-electron DZVP-DFT-orb basis sets for both Sb and O. The total number of basis functions used is 163; the \tilde{X}^2A_1 state of SbO_2 was lowered by 16.77 cm^{-1} through off-diagonal SO interaction.

^dThe computed T_1 diagnostics of the four states shown are 0.0247, 0.0385, 0.0411, and 0.0409, respectively.

^eThe computed T_1 diagnostics of the four states shown are 0.0217, 0.0319, 0.0343, and 0.0338, respectively.

^fAs with footnote a, but the RCCSD(T)/D energies were used for the SO diagonal matrix element; the \tilde{X}^2A_1 state of SbO_2 was lowered by 0.6 cm^{-1} through off-diagonal SO interaction.

← \tilde{X}^2A_1 absorption or LIF band of SbO₂ is expected to be in the visible region at ca. 537 nm (53 kcal/mole or 18 600 cm⁻¹; see later text and Table V), which is significantly lower in energy than the same absorption or LIF band of PO₂ [with T_0 at 326 nm (30 660 cm⁻¹); see Ref. 24, and references therein].

More reliable computed vertical excitation energies (T_{vert} 's) obtained at the MRCI+D and RCCSD(T) levels with larger basis sets are given in Table V. For an excited state of a certain spin symmetry, if its HF wave function converges to the lower state of the same spin symmetry, only CASSCF/MRCI calculations have been carried out. In these CASSCF/MRCI calculations, two roots were requested in order to obtain the energy of the excited state. For an excited state, which has an electronic configuration such that in the SCF procedure of the HF calculations, the configuration remains, RCCSD(T) calculations have also been carried out in addition to CASSCF/MRCI calculations [an example of this is the (2) ²B₁ state; see Table V]. Comparing the calculated results shown in Table V, it is clear that computed CASSCF T_{vert} 's, which lack dynamic electron correlation, are unreliable, as discussed when considering computed T_e 's above. The differences in the computed CASSCF/MRCI+D T_{vert} 's for the electronic states considered, between using basis sets B and C, are less than 1 kcal/mole. The differences in the computed T_{vert} 's between using the CASSCF/MRCI+D and RCCSD(T) methods with the same basis set C are less than 1.5 kcal/mole. The differences in the computed T_{vert} 's between excluding and including Sb 4d¹⁰ core electrons in RCCSD(T) calculations employing basis sets C and D, respectively, are less than 1.5 kcal/mole for all the electronic states considered. These comparisons suggest that the computed T_{vert} values obtained at various higher levels of calculations are reasonably consistent and hence can be concluded to be reliable. The only ambiguous results are the order of the computed T_{vert} 's of the (1) ²A₂ and (1) ²B₂ states, the two lowest excited states of SbO₂. The CASSCF/MRCI+D T_{vert} values with basis sets B and C place the (1) ²A₂ state higher in vertical excitation energy than the (1) ²B₂ state, but the RCCSD(T) values with basis sets C and D have a reverse order. Nevertheless, the differences in the computed T_{vert} values between these two states of SbO₂ are small, particularly at the RCCSD(T) level (less than 0.2 kcal/mole). It should be noted that the computed CI coefficients (larger than 0.77 for the major configuration with basis set C) and T_1 diagnostics (less than 0.035 with basis set D) obtained from the MRCI and RCCSD(T) calculations of the excited states, respectively, for vertical excitation energies from the \tilde{X}^2A_1 state, suggest that multireference character is not important for these excited states. In this connection, it is concluded that the computed RCCSD(T)/D T_{vert} 's are the most reliable with the T_{vert} to the ²A₂ state being lower than to the ²B₂ state by 0.13 kcal/mole (0.33 kcal/mole including spin-orbit coupling; see Table V and later text). Based on the results obtained employing basis sets C and D, the uncertainties associated with the RCCSD(T) T_{vert} 's should be less than 1.5 kcal/mole (0.065 eV). For excited states, where RCCSD(T) T_{vert} 's are not available, the CASSCF/MRCI

+D/C T_{vert} 's are the most reliable. Based on the comparison between RCCSD(T)/D and CASSCF/MRCI+D/C results, the uncertainties associated with the latter values are of the order of 3.0 kcal/mole (0.13 eV).

The computed T_{vert} values including off-diagonal spin-orbit interaction are also shown in Table V. First, results obtained employing the CASSCF/MRCI+D/B energies for the diagonal matrix elements are considered. Those obtained using the BP operator with an all-electron basis set [CASSCF/MRCI+D/B+SO(all-e_8states) in Table V] and spin-orbit pseudopotentials (ECPs) with the ECP basis set ([CASSCF/MRCI+D/B+SO(ecp_4states)]) and [CASSCF/MRCI+D/B+SO(ecp_8states)] are very similar. Also, the results obtained by considering four states (the lowest state of each symmetry) and eight states (two lowest states of each symmetry) are almost identical. Comparing the computed T_{vert} 's between excluding and including spin-orbit coupling, it can be seen that off-diagonal spin-orbit effects on the computed T_{vert} 's are negligibly small except for the (1) ²A₂ and (1) ²B₂ states, which are close in energy as mentioned above. Specifically, the computed T_{vert} values of the (1) ²A₂ state including spin-orbit coupling have increased by ca. 0.57 kcal/mole, when compared with the corresponding unperturbed computed CASSCF/MRCI+D/B T_{vert} values, while those of the ²B₂ state have decreased by ca. 0.25 kcal/mole. With the RCCSD(T)/D energies, which have the (1) ²A₂ state lower in energy than the (1) ²B₂ state, used for the diagonal matrix elements in the average-state CASSCF spin-orbit calculations, the (1) ²A₂ state is lowered by 0.10 kcal/mole by spin-orbit coupling, while the (1) ²B₂ state has increased by 0.10 kcal/mole. The computed lowering in energy of the \tilde{X}^2A_1 state by off-diagonal spin-orbit coupling is negligibly small (less than 0.05 kcal/mole; see footnotes c, d, e, and h of Table V) in all cases. Summing up, it can be concluded that effects of spin-orbit coupling on the computed T_{vert} values of the low-lying electronic states of SbO₂ are negligibly small.

Lastly, a few words on the only available electronic spectrum of SbO₂, the chemiluminescence spectrum of Ref. 26, seem appropriate. SbO₂* may be formed in more than one excited state in the Sb+O₃ reaction, and it is reasonable to assume that the molecular carrier of the chemiluminescence spectrum recorded in the 650–900 nm region is SbO₂ as suggested in Ref. 26. Moreover, it is almost certain that the excited states of SbO₂ formed are vibrationally excited, and it is not unreasonable to assume that the chemiluminescence band recorded is due to emissions to the \tilde{X}^2A_1 state of SbO₂. In this connection, the computed vertical excitation energies from the \tilde{X}^2A_1 state to the excited states of SbO₂ involved in the emissions may be assumed to be approximate upper limits of the high-energy onset of the chemiluminescence band at 650 nm (ca. 44 kcal/mole). Based on the computed vertical excitation energies shown in Table V, the (1) ²B₁ and (2) ²A₁ states with computed T_{vert} values in the 52 kcal/mole (ca. 550 nm) region appear to be suitable candidates for the excited states involved in the chemiluminescence spectrum reported in Ref. 26. Computed transition dipole moments between these two excited states and the \tilde{X}^2A_1 state also

TABLE VI. A summary of some available geometrical parameters (in Å and deg) and vibrational frequencies (bending, symmetrical stretching, and asymmetrical stretching, in cm^{-1}) of the \tilde{X}^2A_1 state of the dioxides of group VA elements, and available T_e values (in kcal/mole) of the \tilde{A}^2B_2 state of these dioxides.

\tilde{X}^2A_1	Method	r_e	θ_e	$\omega_e(a_1; a_1; b_2)$	$T_e(\tilde{A}^2B_2)$
NO ₂	MW ^a	1.1934	134.1	758.636; 1347.53; 1669.02	27.9
	LIF/DF ^b				
PO ₂	LMR/MW; ^c r_0 , average ω	1.4665	135.3	377; 1090; 1278	14.45
	IR laser ^d			⋯; ⋯ 1327.53452(69)	
	LIF/DF ^e			397.3(43); 1075.4(50); ⋯	
	MRDCI/TZ+2d+R ^f	1.464	135.14	389; 1052; 1338	
	MP2/6-311G(2d) ^g	1.478	136.6	403; 1085; 1486	
	RCCSD(T)/aug-cc-pVQZ ^g	1.474	134.8	384; 1069; ⋯	
	RCCSD(T)/aug-cc-pVQZ ^h	1.474	134.8	390; 1073; 1349	
AsO ₂	MP2/[6s4p2d], [4s2p1d] ⁱ	1.630	135		
SbO ₂	RCCSD(T)/B	1.8363	122.81	205; 762; 829	5.0
	RCCSD(T)/E	1.8271	121.75		4.38
	CBS				4.1

^aReferences 23 and 46.

^bReferences 37 and 38.

^cReference 47.

^dReference 48.

^eReference 45.

^fReference 41.

^gReference 24.

^hReference 49.

ⁱReference 34.

suggest that emissions from the former states to the \tilde{X}^2A_1 state should have significant intensities (see Table IV). In conclusion, it is not inconsistent, based on *ab initio* results obtained in the present study, that the chemiluminescence spectrum recorded in the 650–900 nm region is due to SbO₂^{*}. However, further spectroscopic and computational studies will be required to firmly identify the molecular carrier of, and the electronic and vibrational states involved in, the chemiluminescence spectrum recorded in the 650–900 nm region resulting from the reaction between Sb and O₃ in the gas phase.²⁶

CONCLUDING REMARKS

DFT and high-level *ab initio* calculations have been carried out on some low-lying electronic states of SbO₂. These calculations have clarified the uncertainty associated with the geometrical structure of the ground electronic state of SbO₂ reported previously.^{16,22} The ground state of SbO₂ has been unambiguously established to be the \tilde{X}^2A_1 state with a C_{2v} structure, similar to the dioxides of other lighter group 15 elements. In addition, a number of low-lying electronic states of SbO₂ have been characterized. Among them, it should be noted that the \tilde{A}^2B_2 state of SbO₂ has a best computed T_e of only 4.1 kcal/mole (1434 cm^{-1} , at the CBS limit). The closeness in energy of the \tilde{A}^2B_2 state to the \tilde{X}^2A_1 state will contribute to the thermodynamic properties of SbO₂.

Reliable vertical excitation energies from the \tilde{X}^2A_1 state to the low-lying electronic states of SbO₂ have been computed. In addition, transition dipole moments (TDMs) between the \tilde{X}^2A_1 state and low-lying excited states of SbO₂ have been calculated. These computed TDMs suggest that

the strongest electronic transition between the \tilde{X}^2A_1 state of SbO₂ and low-lying excited states is the (2) $^2A_1 \leftarrow \tilde{X}^2A_1$ transition, similar to a conclusion made in our previous combined *ab initio*/Franck-Condon study on the LIF and single vibronic level emission spectra of PO₂.²⁴ However, the computed vertical excitation energy of this transition of SbO₂ obtained in the present study suggests that the absorption/LIF band of this transition should be in the visible region, significantly lower in energy than the same transition of PO₂, which is in the ultraviolet region. The calculated results reported here will be useful to assist interpretation of the absorption and/or LIF spectra of SbO₂ which are/is yet to be recorded.

A summary of some available geometrical parameters and vibrational frequencies of the \tilde{X}^2A_1 state of the dioxides of the group 15 (or VA) elements and available T_e values of the \tilde{A}^2B_2 state of these dioxides are given in Table VI. To our knowledge, there are no such data, theoretical or experimental, available for BiO₂, the heaviest dioxide of the series. For AsO₂, the optimized geometrical parameters of the \tilde{X}^2A_1 state have been reported at the MP2 level using the [6s4p2d] and [4s2p1d] basis sets for As and O, respectively.³⁴ In addition, a recent computational study on cyclic AsO₂ and AsS₂ systems has reported optimized geometrical parameters and harmonic vibrational frequencies of the \tilde{A}^2B_2 state, and also other cyclic excited states of AsO₂ obtained employing the MP2 and DFT (with different functionals) methods.²⁵ However, although the relative energies of these cyclic states were calculated at higher levels of theory than the MP2 level, and a lower 2A_1 state, which corresponds to the \tilde{X}^2A_1 state of AsO₂ reported in Ref. 34, was also mentioned in Ref. 25, the

relative energy of the \tilde{A}^2B_2 state with respect to the \tilde{X}^2A_1 state of AsO₂ has not been reported in Ref. 25. Consequently, the T_e value of the \tilde{A}^2B_2 states of AsO₂ is unavailable. The authors of Ref. 25 have considered the \tilde{A}^2B_2 state of AsO₂ as the ground state of a cyclic AsO₂ rather than as an excited state of AsO₂. For NO₂ and PO₂, both theoretical and experimental geometrical parameters and vibrational frequencies are available (see Table VI). For the T_e of the \tilde{A}^2B_2 state of NO₂, although there is significant vibronic interaction between the \tilde{X}^2A_1 and \tilde{A}^2B_2 states near the conical intersection of the two electronic surfaces resulting in chaotic behavior being observed in the 10 000 cm⁻¹ region of the absorption and LIF spectra of NO₂ (see Refs. 35 and 36, and references therein), the T_e value of the \tilde{A}^2B_2 state has been established to be 9760 ± 100 cm⁻¹,³⁷ [27.9 kcal/mole; T_0 = 9734 cm⁻¹ (Ref. 38)]. For the T_e of the \tilde{A}^2B_2 state of PO₂, however, only theoretical values are available (see Table VI).

Comparing the available equilibrium geometrical parameters of the \tilde{X}^2A_1 state of the dioxides of the group 15 elements, it is noted that the bond angles for NO₂, PO₂, and AsO₂ have very similar values, ranging between 134° and 135°, but SbO₂ has a significantly smaller bond angle of 121.8° (see Table VI). This may be compared with some group 16 (VIA) analog, SO₂, SeO₂, and TeO₂, which have bond angles of 119.5°,²³ 113.83°,³⁹ and 111.15° [RCCSD(T)/ECP46MWB-aug-cc-pV5Z, aug-cc-pV5Z value from Ref. 40], respectively. In the case of the dioxides of the group 16 elements, the significant decrease of the bond angle occurs between SO₂ and SeO₂. The smaller magnitude of the decrease in the bond angle between SO₂ and SeO₂ (ca. 6°) than that between AsO₂ and SbO₂ (ca. 13°) is almost certainly due to the extra valence electron in the group 16 elements and the already smaller angle in the lighter members of the group 16 dioxides than those of group 15 dioxides. In this connection, it will be interesting to see how far the bond angle would decrease with BiO₂.

Comparing available vibrational frequencies of the dioxides of the group 15 elements, as shown in Table VI, the symmetric stretching frequency is smaller than the asymmetric stretching frequency for the \tilde{X}^2A_1 state of both NO₂ and PO₂. The computed RCCSD(T)/B vibrational frequencies of SbO₂ reported here (see Table II) behave in the same way as the lighter members of the series with the symmetric stretching frequency being smaller than the asymmetric stretching frequency. This comparison suggests that the pattern is the same over the whole series. In this connection, the computed DFT vibrational frequencies of SbO₂, which give larger values for the symmetric stretching mode than the asymmetric stretching mode (see Table II), are almost certainly unreliable, as concluded above.

Another point of interest from the comparison of the data shown in Table VI is the trend in the separations between the \tilde{A}^2B_2 and \tilde{X}^2A_1 states, i.e., T_e of the \tilde{A}^2B_2 state of the dioxides of the group 15 elements. Although the T_e value of the \tilde{A}^2B_2 state for AsO₂ is not available, it can be seen that the separations between the \tilde{A}^2B_2 and \tilde{X}^2A_1 states decrease from NO₂ to SbO₂. In this connection, it seems clear

that the MP2/6-311+G(2d) $T_e(\tilde{A}^2B_2)$ value of 32.0 kcal/mole from Ref. 24 for PO₂ is almost certainly not as reliable as the MRDCI/TZ+2d+R value of 14.45 kcal/mole.⁴¹ (It is noted that the UHF wave function of the \tilde{A}^2B_2 state of PO₂ used in the MP2 calculation of Ref. 24 has a computed $\langle S^2 \rangle$ value of 0.809, suggesting that spin contamination from higher-spin quartet states may have affected the MP2 results.) Following the trend, as shown in Table VI, it is expected that the $T_e(\tilde{A}^2B_2)$ value of AsO₂ should lie between 14.45 and 4.1 kcal/mole. In addition, the $T_e(\tilde{A}^2B_2)$ value of BiO₂ should be smaller than 4 kcal/mole, and there is a possibility that for BiO₂ the ground state may be a 2B_2 state rather than a 2A_1 state.

Regarding the appropriate computational methods to be used in studying the low-lying electronic states of SbO₂ and related systems, such as AsO₂ and BiO₂, the results obtained from the present study shows that both the DFT (with some most commonly used functionals) and MP2 methods are inadequate at least in some aspects. Higher level methods, such as the RCCSD(T) and/or CASSCF/MRCI methods, should be employed in order to obtain reliable results. Regarding the requirements on the basis set to be used, a large core ECP, such as the ECP48MWB, with a quadruple-zeta quality valence basis set for Sb has been shown to be reasonably adequate. In addition, contributions from both outer core correlation and spin-orbit coupling have been found to be small.

Lastly, preliminary calculations on low-lying quartet states of SbO₂ have been carried out and they suggest that the lowest quartet state is a 4A_1 state, which is computed to be ca. 54 kcal/mole above the \tilde{X}^2A_1 state at the CASSCF/MRCI+D/B level of calculation. It should be noted that spin-orbit interaction between doublet and quartet states can be significant, if the states involved are close in energy. However, in relation to the few lowest-lying doublet states considered in the present study, low-lying quartet states of SbO₂, which are considerably higher in energy, should not affect the energy positions of these doublet states significantly via spin-orbit interaction. A full report of the investigation on low-lying quartet states of SbO₂ and also low-lying singlet and triplet states of SbO₂⁺ is forthcoming. On the experimental front, we call for spectroscopic studies on SbO₂ (and/or similar systems, such as AsO₂ and BiO₂) to be carried out, for example, by gas phase supersonic jet-cooled LIF spectroscopy. SbO₂ may be prepared in the gas phase for spectroscopic study by pyrolysis (as used in the LIF studies of TeO₂ (Ref. 42) and SeO₂,⁴³ though a multi-stage oven would probably be required) or laser ablation (see, for example, Ref. 44) of solid antimony oxide, photolysis or electric discharge of some suitable precursors [e.g., SbCl₃ or SbCl₅+O₂; the photolysis of a mixture of PCl₃ and O₂ molecules was employed in the LIF study of PO₂ (Ref. 45)], or some suitable reactions between Sb atoms or small clusters (produced by laser ablation or pyrolysis) and an oxidizing agent (e.g., O atom produced in a microwave discharge or O₃ in the throat of a supersonic nozzle).

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