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Experimental and theoretical study of the Sn–O bond formation between atomic tin and molecular oxygen†

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The merging of the electronic structure calculations and crossed beam experiments expose the reaction dynamics in the tin (Sn, 3P_j) – molecular oxygen (O₂, X³ Σ_g^-) system yielding tin monoxide (SnO, X¹ Σ^+) along with ground state atomic oxygen $O(^3P)$. The reaction can be initiated on the triplet and singlet surfaces via addition of tin to the oxygen atom leading to linear, bent, and/or triangular reaction intermediates. On both the triplet and singlet surfaces, formation of the tin dioxide structure is required prior to unimolecular decomposition to SnO($X^1\Sigma^+$) and O(3P). Intersystem crossing (ISC) plays a critical role in the reaction mechanism and extensively interosculates singlet and triplet surfaces. The studied reaction follows a mechanism parallel to that for the gas phase reaction of germanium and silicon with molecular oxygen, however, the presence of the tin atom enhances and expands ISC via the "heavy atom effect".

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Introduction

Since the exploitation of the main group XIV element tin (Sn) merged with the transition metal copper (Cu) in bronze alloys more than 5000 years ago,¹ the preparation and properties of subvalent tin (II) compounds together with their isovalent carbon (C), silicon (Si), and germanium (Ge) counterparts have intrigued the computational chemistry, physical inorganic, and preparative synthetic communities from the perspectives of electronic structure theory and chemical bonding.² From carbon to tin, the electronic structure of the dihydrides transforms noticeably. Whereas for the lightest main group XIV element carbon, the divalent hydride methylene (CH_2, X^3B_1) has a triplet electronic ground state and a singlet–triplet splitting to the a $^1\mathrm{A}_1$ state of 36 to 38 kJ mol $^{-1}$,^{3,4} all higher homologues silylene (SiH₂), germylene (GeH₂), and stannylene (SnH₂) hold an X^1A_1 electronic ground state. The singlet-triplet splitting to the a ${}^{3}B_{1}$ state increases from 80–88 kJ mol $^{-1}$ in silylene $(SiH₂)^{5–11}$ *via* 100–105 kJ mol $^{-1}$ in

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germylene $(\text{GeH}_2)^{12}$ and 98–111 kJ mol $^{-1}$ (ref. 12,13) in stannylene (SnH2) as the period increases. This trend can be rationalized in terms of discrete sizes of the valence orbitals of carbon, silicon, germanium, and tin; the enhanced size of the valence orbitals of silicon, germanium, and tin compared to carbon results in an ineffective hybridization of the s and p orbitals in silicon, germanium, and tin; this is accompanied by the stabilization of the singlet versus triplet state in silylene, germylene, and stannylene. This finding correlates nicely with the reduction of the H–E–H angles ($E = C$, Si, Ge, Sn) in the electronic ground states, which are reduced from 129.8° (CH₂, X^3B_1) via 93.4° (SiH₂, X^1A_1) to 92.3° (GeH₂, X^1A_1) and 92.4° (SnH₂, X^1A_1).^{14,15} The diminished reactivity of stannylene $(SnH₂)¹⁶$ compared to their isovalent methylene (CH_2) ,^{17–21} silylene (SiH_2) ,^{22–24} and germylene (GeH₂) analogues²⁵ culminated in the preparative synthesis of subvalent tin (n) compounds such as $\text{Sn}_2[(\text{Me}_3\text{Si})_2\text{CH}]^{26}$ and $\text{Sn}[\text{N}(\text{SiMe}_3)]_2$.²⁷

The mounting attention in the (in)organic tin(π) chemistry and industrial applications of $tin(n)$ oxide (SnO) in conjunction with indium tin oxide (ITO) on touchscreens and thin film transistors $(TFT)^{28,29}$ revitalized widespread research in the electronic structure and chemical bonding of binary oxides of main group XIV elements. In gas-phase carbon monoxide (CO), two π and one σ -bond generate a carbon–oxygen triple bond. $30,31$ Whereas carbon monoxide is a gas at 293 K, gasphase silicon monoxide (SiO), is unstable and disproportionates to amorphous silicon and silicon dioxide $(SiO₂)$ ^{32,33} In the diatomic oxides, the bond lengths increase from carbon via silicon and germanium to tin from 112.8 pm via 151.2 pm and 161.7 pm to 183.3 pm. 34 The electronegativity difference

[†] Electronic supplementary information (ESI) available: Potential energy surface with included bond distances (Å) and bond angles of each transition state, intermediate, minimum on the seam of crossing. Optimized Cartesian coordinates and vibrational frequencies for all intermediates, transition states, MSX, reactants, and products involved in the reactions of the atomic tin $(Sn; {}^{3}P_{j})$ with oxygen $(O_2; X^3\Sigma^-_g)$. See DOI: <https://doi.org/10.1039/d4cp03687e>

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between the main group XIV element and oxygen rises from 1.00 in carbon monoxide (CO) to 1.78 in tin monoxide (SnO) on the Pauling scale suggesting an enhanced ionic character in the tin–oxygen bond compared to the carbon–oxygen bond. Also, within main group XIV, the +II oxidation state becomes increasingly stable as demonstrated by the stability of commercially available tin(π) and tin(π) compounds such as tin(π) chloride and tin($|v\rangle$ sulfide. Nevertheless, tin($|v\rangle$) is still considered to be more stable than $tin(n)$; as such, tin monoxide (SnO) emerged as a reduction agent that can be oxidized easily to tin dioxide $(SnO₂)$ ³⁵ Chlorides and organic compounds such as tin tetrachloride (SnCl₄), tetramethyltin ((CH₃)₄Sn), and dimethyldichlorotin $((CH₃)₂SnCl₂)$ are used as precursors for the gasphase synthesis of tin dioxide. These precursors decompose³⁶⁻³⁸ into atomic tin, which also makes studying the reaction of tin plus molecular oxygen (O_2) an indispensable step to untangle the reaction mechanism during the synthesis of tin dioxide. Previously, the kinetics of the reaction of atomic tin with molecular oxygen was studied at high temperatures of 1300 to 2600 K using a shock tube equipped for atomic resonance absorption spectroscopy $(ARAS)$,^{39,40} in a fast-flow reactor with laser-induced fluorescence (LIF) over the wide temperature range 380 to 1840 K., 41 and by flash photolysis absorption spectroscopy at 295 K. $42-44$ Studies demonstrate that the reaction is spin-allowed and the main products are tin monoxide (SnO, $X^1\Sigma^+$) and atomic $O(^{3}P)$. Rate constants measurements show that the reaction is relatively fast, with a slight positive temperature dependence (Table 1). It must be mentioned that in $1978⁴⁵$ the reaction of atomic tin with molecular oxygen was tried in a crossed molecular beam setup with a high-temperature (1525 K) molten source for the tin beam. However, the authors were unable to experimentally verify the reaction energy, and the level of development of quantum chemical calculations at the moment would not have allowed them to elucidate the reaction mechanism of such an electron-rich system.

However, whereas tin dioxide $(SnO₂)$ has been successfully synthesized in the gas phase exploiting flame spray pyrolysis and droplet combustion techniques,⁴⁶ no route has been developed to prepare divalent tin monoxide (SnO) in a directed synthesis. Here, we report the gas-phase preparation of tin monoxide (SnO, $(X^1\Sigma^+)$ through the bimolecular reaction of ground state tin atoms (Sn; ³P_j) with molecular oxygen (O₂; $X^3\Sigma_g^-$) utilizing the crossed molecular beams technique.^{47,48} An elucidation of elementary reactions at the microscopic level in tandem with electronic structure calculations affords remarkable insights into the intimate reaction dynamics through which highly reactive subvalent oxides like tin oxide (SnO, $X^1\Sigma^+$) can be generated via a single collision event between atomic tin and a prototype oxidant $(O_2; X^3\Sigma_g^-)$ under single-collision conditions in the gas phase through non-adiabatic reaction dynamics. This reaction is also of fundamental importance to the reaction dynamics community as a benchmark of triatomic systems involving the 'heavy' main group XIV element tin, which has been elusive until now, and the comparison with the isovalent $Si-O_2^{49}$ and $Ge-O_2^{50}$ systems explored previously.

Experimental and computational

Experimental

The gas-phase reaction of atomic tin $(Sn, \, {}^3P_j)$ with molecular oxygen $(O_2, X^3\Sigma_g^-)$ was carried out under single-collision conditions using the crossed molecular beams machine.^{47,51} The experimental setup, data acquisition, and data processing are described elsewhere in detail.⁴⁷⁻⁴⁹ Briefly, the reactant atomic tin $(Sn, {}^{3}P_{j})$ was generated *in situ* in the primary source chamber by laser ablation (3 mJ, 266 nm, 30 Hz; Quanta-Ray) from a rotating tin rod (99.98% Sn, Alfa Aesar) and seeding the ablated atoms in a pulsed argon beam (Ar, 99.9999%; Airgas) that was released from a piezoelectric valve operating at 60 Hz and a backing pressure of 4 atm (Fig. 1a). All five most abundant isotopes of tin^{52} were detected in the beam with the signal ratios matching their natural abundance (Table 2). The atomic tin beam was skimmed and velocity selected using a four-slot chopper wheel; this achieved a peak velocity $v_p = 631 \pm 22$ m s⁻¹ and speed ratio $S = 8.5 \pm 0.6$. Neat oxygen gas (550 Torr, O_2 , 99.998%; Matheson) was used as a precursor to produce the 'reactant two' supersonic molecular oxygen beam (60 Hz, $v_{\rm p}$ = 776 \pm 20 m s⁻¹, *S* = 15.6 \pm 1.0) in the secondary source chamber. Both molecular beams intersected at an angle of 90° in the scattering chamber at a mean collision energy of $E_C = 13 \pm 1$ kJ mol⁻¹. The reactively scattered products were ionized by electron ionization at 80 eV (2 mA) at the entrance of the rotatable detector, filtered according to m/z by the QMS (Extrel, QC 150; 1.2 MHz), and detected using a Daly-type particle ion counter (Fig. S1, ESI†). 53 Angularly resolved time-of-flight (TOF) spectra were recorded at discrete laboratory angles in 2.5° steps (Fig. 1b). Operating the laser at half frequency of the pulsed valves and sufficient length of the recorded TOFs allowed an

Table 1 Experimentally measured rate constants for reaction of Sn(³P_j) with O₂

Method ^a	k, cm ³ molecule ⁻¹ s ⁻¹	Temperature range, K	Source
Shok tube, ARAS	$10^{-9.80}\times {\rm e}^{-11.7\,{\rm kJ\,mol}^{-1}/{\rm RT}}$	1350–2600	Ref. 39
	$10^{-9.41 \pm 0.03} \times e^{-11.5 \pm 1.1 \text{ kJ mol}^{-1} / RT}$	1300-2250	Ref. 40
Fast-flow reactor, LIF	$10^{-12.29} (T/K)^{0.79} \times e^{-3.6 \text{ kJ} \text{ mol}^{-1} / \text{RT}}$	380-1840	Ref. 41
Flash photolysis absorption	$1.05 \pm 0.08 \times 10^{-11}$	295	Ref. 42
	$Sn(^{3}P_{0})$: 2.1 \pm 0.1 \times 10 ⁻¹¹		Ref. 43
	$\rm Sn(^3P_0): 3.5\pm0.4 \times 10^{-11}$		Ref. 44
	$\rm Sn(^3P_1): 8.2\pm0.5 \times 10^{-11}$		

 $\text{Sn}({}^{3}P_{2})$: 4.91 \pm 0.3 \times 10⁻¹¹

 a^a ARAS – atomic resonance absorption spectroscopy, LIF – laser-induced fluorescence.

Fig. 1 Experimental setup. (a) Side view on the cross point; (b) Top-view of the experiment with the detector.

Table 2 Theoretical and experimental ratios of signal for different tin isotopes in the received atomic tin beam in argon

	120 Sn	$^{119}\mathrm{Sn}$	118 Sn	117Sn	$^{116}\mathrm{Sn}$
Abundance	32.6%	8.6%	24.2%	7.7%	14.5%
Theoretical ratio of signal ^{<i>a</i>}		0.26	0.75	0.24	0.44
Experimental ratio in beam		0.25 ± 0.01	0.74 ± 0.01	0.24 ± 0.01	0.49 ± 0.01
a Ratio to the most abundant isotope 120 Sn.					

instant background subtraction (''laser-on'' minus ''laser-off'') during the TOF recording.

To gain information on the reaction dynamics, TOF spectra and the laboratory angular distribution (LAD) were transformed from the laboratory to the center-of-mass frame by a forwardconvolution routine.^{54,55} This approach uses initially a trial angular flux $T(\theta)$ and translational energy $P(E_T)$ distributions in the center-of-mass (CM) frame to simulate the TOFs and LAD (laboratory data). CM functions were iteratively varied until the best fit of the TOF spectra and LAD was achieved. Together the CM functions constitute the reactive differential cross sections $I(\theta, u) \sim$ $P(u) \times T(\theta)$, where u – the CM velocity, θ – angle in the CM system. The reactive differential cross sections can be represented as a flux contour map that depicts the probability of the products to scatter at the specific angle (θ) with the specific kinetic energy (u) .

Computational

Being a heavy atom with 50 electrons, tin requires consideration of relativistic effects by using a relativistic effective core potential (ECP).⁵⁶⁻⁵⁸ 28 electrons at the $1s^22s^22p^63s^23p^63d^{10}$ set of orbitals of Sn are described by means of Stuttgart ECP28MDF⁵⁷ and therefore, the number of electrons on the tin atom explicitly included in the computations is 22 at the $4s^24p^65s^24d^{10}5p^2$ set of orbitals, making the total number of electrons for the $SnO₂$ system to be 38. The hybrid ω B97X-D density functional theory⁵⁹ with Dunning's augmented correlation-consistent triple- ζ basis sets aug-cc-pVTZ (for O) and aug-cc-pVTZ-PP (for Sn) $56,57,60$ in the Gaussian 09 software package 61 was originally applied to optimize geometries and to compute vibrational frequencies of the intermediates and transition states on the $SnO₂$ potential energy surface (PES). All the structures were subsequently

reoptimized in Molpro 2015^{62} utilizing the multireference second-order perturbation theory (CASPT2) method $63,64$ with the augmented quadruple- ζ basis sets aug-cc-pVOZ (for O) and aug-cc $pVQZ-PP$ (for Sn)^{56,57,60} and with full valence active space containing 16 electrons distributed on 12 orbitals $(5s^25p^2$ for tin and two sets of $2s^22p^4$ for oxygen). The CASPT2 wavefunctions used initial orbitals taken from full valence, multireference complete active space self-consistent field (CASSCF) calculations^{65,66} with the same quadruple- ζ basis sets.^{56,57,60} All calculations were carried out within the C_s point group, with the numbers of orbitals included in the active space ('occupied') and those kept closed being $17A' + 6A''$ and $8A' + 3A''$, respectively, thus resulting in 12 active orbitals $(9A' + 3A'')$. It should be noted that in the CASPT2 calculations $4s^24p^6$ electrons of tin and $1s^2$ of each oxygen were treated as a core $(5A' + 1A'')$ and not included in single and double excitations, with only $4d^{10}$ (3A' + 2A") electrons of Sn involved beyond the active space. Thus, 26 electrons of $SnO₂$ participated in the dynamic electronic correlation calculations.

The minimal energy structures on the seams of crossing (MSX) between singlet $({}^{1}A')$ and triplet $({}^{3}A'$ or ${}^{3}A'')$ states and between two different triplet states $(1^3A' - 2^3A'$ or $A' - A''$ were located using Molpro 2015 using two alternative approaches. First, the CASSCF(16,12) method with the Karlsruhe segmented contracted, split-valence triple- ζ def2-TZVPPD basis set containing two sets of polarization functions and a set of diffuse function^{67,68} was employed for the conical intersection optimization followed by CASPT2(16,12)/aug-cc-pVQZ-(PP) single-point calculations for the MSX energy refinement. Where the direct MSX optimization was unsuccessful, we utilized CASSCF(16,12)/aug-cc-pVQZ-(PP) (PP relates only to Sn) and CASPT2(16,12)/aug-cc-pVQZ-(PP) single-point computations to carry out two-dimensional potential

energy scans separately for each state in relevant regions of the PES. Such regions were identified based on the constructed potential energy diagram for the electronic states involved.

Furthermore, the energies of the reactants and products were calculated both at the coupled clusters⁶⁹ CCSD(T)/CBS// oB97X-D/aug-cc-pVTZ-(PP) level with the complete basis set (CBS) extrapolation from the values obtained with the aug-ccpVOZ- (PP) and aug-cc-pVTZ- (PP) basis sets for isolated Sn, O_2 , SnO, and O and at the CASPT2(16,12)/aug-cc-pVQZ-(PP)/ ω B97X-D/aug-cc-pVTZ-(PP) level considering the reactants $Sn + O₂$ and the products $SnO + O$ as supermolecules, with two fragments positioned sufficiently far from one another so that their interaction is negligible. The two theoretical methods used have resulted in practically the same reaction energy. Finally, potential energy scans at the CASPT2(16,12)/aug-cc-pVQZ-(PP) level of theory were implemented to explore the entrance O_2 addition channel for the formation of an initial triplet intermediate and the exit O elimination channel on the triplet PES and to verify barrierless connections between the $SnO₂$ intermediates with reactants/products.

Results and discussion

Laboratory frame

The ratios of the natural isotope abundances of tin 116 Sn, 117 Sn, 118 Sn, 119 Sn, 120 Sn, 122 Sn, and 124 Sn are 14.5, 7.7, 24.2, 8.6, 32.6, 4.6 and 5.8%, respectively. Therefore, the reactive scattering signal for tin monoxide (SnO) was initially monitored in the range of m/z 132–141. The ratio of the collected signal of tin monoxide at the center-of-mass angle nicely matches the prediction from the natural abundance of tin isotopes (Fig. 2). Non-zero signals at $m/z = 137$ and 139 arise from the presence of 18 O isotopes.

These raw data alone demonstrate a single reaction channel via the emission of atomic oxygen (16 amu) and formation of tin monoxide $(^{116}Sn^{16}O^+, m/z = 132; ^{117}Sn^{16}O^+, m/z = 133; ^{118}Sn^{16}O^+,$ $m/z = 134;$ $^{119}Sn^{16}O^+, m/z = 135;$ $^{120}Sn^{16}O^+, m/z = 136;$ $^{122}Sn^{16}O^+, m/z$ $z = 138$; 124 Sn¹⁶O⁺, $m/z = 140$) in the reaction of atomic tin with molecular oxygen. The corresponding TOF spectra were collected at the best signal-to-noise ratio at $m/z = 136$ (Fig. 3) and were then normalized to the signal at the CM angle to obtain the laboratory angular distribution (LAD). More than 2.5 \times 10⁶ TOF spectra were recorded between 8.75[°] $\leq \theta \leq 58.75$ [°] with the tin beam defined as $\Theta = 0^{\circ}$ direction. The LAD is rather wide, starts at 53.75°, and extends beyond the range of the rotating detector at small angles (less than 8.75°). The TOF spectra are broad from at least 700 to 1500 µs and peak between 750 and 900 µs. These findings propose that the tin monoxide products were formed via indirect scattering dynamics through complex formation involving $SnO₂$ intermediates.47,70,71

Center-of-mass frame

To further illuminate the underlying reaction mechanism(s) accompanied by the potential involvement of intersystem crossing (ISC), excited state surfaces, and non-adiabatic reaction

Fig. 2 The ratio of the signal for the reaction of atomic tin with molecular oxygen at the center-of-mass angle at different m/z. The blue bars are the experimental results, red – predicted ratio according to the natural isotope abundance of tin.

dynamics, a transformation of the laboratory data from the laboratory reference frame into the center-of-mass reference frame is accomplished using a single channel fit of the laboratory data and reaction (1):

$$
{}^{120}\mathrm{Sn}({}^{3}\mathrm{P}_{j})+{}^{16}\mathrm{O}_{2}(X^{3}\Sigma_{g}^{-})\,\to\,{}^{120}\mathrm{Sn}^{16}\mathrm{O}(X^{1}\Sigma^{+})+{}^{16}\mathrm{O}({}^{3}\mathrm{P}). \quad \ \ (1)
$$

The best-fit CM functions are depicted in Fig. 4a and b. The error ranges of the $P(E_T)$ and $T(\theta)$ functions are determined within the 1σ limits of the corresponding laboratory angular distribution and beam parameters (beam spreads and beam velocities) while maintaining a good fit of the laboratory TOF spectra and LAD. The translational energy flux distribution $P(E_T)$ (Fig. 4a) contains valuable information about reaction dynamics and thermodynamics. The derived $P(E_T)$ distributions exhibit a maximum translational energy release (E_{max}) of 76 \pm 12 kJ mol⁻¹. Energy conservation dictates that for those molecules born without internal excitation, E_{max} is the sum of the collision energy (E_C) and the reaction energy. Taking into account the collision energy of 13 \pm 1 kJ mol⁻¹, the reaction energy was determined to be exoergic by 63 \pm 13 kJ mol⁻¹. Thermodynamic analysis of our experimental results proves the observation of reaction (1) with stated electronic and spin–orbit levels. In the case of the presence of only the $Sn(^{3}P_{0})$ in the beam reaction energy would be -24 kJ mol⁻¹, while reaction energy calculated for the $Sn(^{3}P_{2})$ provides a close match $(-65 \text{ kJ mol}^{-1})$ with the experimental data. Sn(${}^{3}P_{0}$) and $Sn(^{3}P_{1})$ are also present in the beam but masked in the lower energy section of the CM translational energy distribution (Fig. 4a). Production of higher energetic species $\text{SnO}(a^3\Sigma^+)$ or $O(^{1}D)$ drives the reaction endoergic to +208⁷² or +166 kJ mol⁻¹, respectively. These reactions are closed under our experimental

Fig. 3 (a) Laboratory angular distribution (b) and time-of-flight (TOF) spectra recorded at $m/z = 136$ for the reaction of the tin atom with molecular oxygen at a collision energy of 13 \pm 1 kJ mol $^{-1}$. The circles represent the experimental data and the solid lines are the best fits, the green atom is tin, red atoms are oxygen.

conditions with E_{C} = 13 \pm 1 kJ mol $^{-1}$. The average translational energy of the products was derived to be 28 \pm 5 kJ mol $^{-1}$, suggesting that 38 \pm 12% of the total available energy is channeled into the translational degrees of freedom of the products, which usually implies that the reaction mechanism proceeds through the formation of a covalently bound intermediate.47,70,71 The center-of-mass angular distribution $T(\theta)$ can provide additional information about the reaction dynamics (Fig. 4b). The 'flat' $T(\theta)$ reveals that products after a collision are scattered in all directions with equal probability (isotropic scattering). The forward-backward symmetry and intensity distribution also propose indirect scattering dynamics *via* long-lived (longer than their rotational periods) $SnO₂$ complex(es).⁷³ Foresaid findings are also enclosed in the flux contour map (Fig. 4c), which depicts an overall image of the reaction scattering process.

Reaction mechanism

Now we combine our experimental results with electronic structure calculations to unlock the underlying chemical dynamics and reaction mechanism(s) of atomic tin oxidation in the gas phase. For the Sn(${}^{3}P_{j}$) – O₂(X ${}^{3}\Sigma_{g}^{-}$) reaction, both reactants are in their triplet electronic ground states; since the

Fig. 4 (a) Center-of-mass translational energy $P(E_T)$, (b) angular $T(\theta)$ flux distributions, and (c) flux contour map for the reaction of the tin atoms with molecular oxygen. The solid lines represent the best fit, while the shaded areas indicate the error limits. For $T(\theta)$, the direction of the tin beam is defined as 0° and of the oxygen as 180 $^{\circ}$. The green atom is tin, red atoms are oxygen.

accessible products are tin monoxide (SnO, $X^1\Sigma^+$) and atomic oxygen $O(^{1}D/^{3}P)$, close scrutiny has to be attributed to the triplet and singlet $SnO₂$ surfaces. Here, the computations identified three singlet (i1, i4, and i8) and five triplet (i2, i3, i5, i6, $i7$) SnO₂ intermediates, six transition states (ts1-ts6), five singlet–triplet seams of crossings (MSX1a, MSX1b, MSX2, MSX7, and MSX8), and eight triplet–triplet seams of crossings (MSX3–MSX6, MSX9–MSX12).

The singlet surface (Fig. 5, red) can be accessed via two barrierless entrance channels: end-on addition with the formation of a linear (SnOO, i4, C_{∞} v, $^1\Sigma^+$) or side-on addition leading to a triangular (SnO₂, i1, C_{2v} , $^{1}A_{1}$) intermediate. The triangular intermediate i1 can undergo either Sn–O bond

Fig. 5 Potential energy surface (PES) of the reaction of atomic tin (Sn, ³P_i) with oxygen (O₂, X³∑₅). The italic numbers colored red, blue, black, and pink give the energies at the CASPT2(16,12)/aug-cc-pVQZ-(PP) (PP relates to Sn) level of theory with ZPE at the oB97X-D/aug-cc-pVTZ-(PP) level of theory. The reaction energies of the products are calculated using CCSD(T)/CBS(aug-cc-pV(T+Q)Z-(PP))//oB97X-D/aug-cc-pVTZ-(PP). The geometries of MSXs are either optimized at the CASSCF(16,12)/def2-TZVPPD level of theory, with their single-point energies recalculated at the CASPT2(16,12)/aug-ccpVQZ-(PP) level of theory, or located using two-dimensional scans of the PES at the CASPT2(16,12)/aug-cc-pVQZ-(PP) level of theory. The energies are shown in kJ mol⁻¹. The tin atoms are colored green, and the oxygen atoms are colored red. Detailed structural parameters of the intermediates, transition states, and MSX are given in Fig. S2 in ESI.†

rupture to $i4$ *via* a high barrier of 201 kJ mol $^{-1}$ or O–O bond cleavage via a 99 kJ mol $^{-1}$ barrier to the linear tin dioxide structure ${\bf i}$ 8 (OSnO, $D_{\infty \text{h}},$ $^1\Sigma^+_g$). Both ${\bf i}$ 4 and ${\bf i}$ 8 have an only spinallowed exit channel without a transition state to tin monoxide $\text{SnO}(\text{X}^1 \Sigma^{+})$ plus O(^1D) with a total reaction endothermicity of 125-166 kJ mol $^{-1}$ (depending on the initial state of the Sn atom). On the triplet surface (Fig. 5, blue), the reaction can be initiated via the addition of ground-state tin to one of the oxygen atoms of molecular oxygen leading to the bent triplet intermediates **i3** (\angle SnOO = 103°, C_s , ³A″) or **i5** (\angle SnOO = 121°, C_s , 3 A') via tiny barriers of 1 or 3 kJ mol⁻¹, respectively. Intermediates i5 and i3 can isomerize via migration of the terminal oxygen atom to the tin atom accessing bent tin dioxide structures **i6** (\angle OSnO = 101[°], C_{2v} , ³B₂) and **i7** (\angle OSnO = 68[°], C_{2v} , ${}^{3}B_{2}$), respectively. Both i6 and i7 undergo a barrierless, unimolecular decomposition to tin monoxide (SnO, C_{∞} v, $\mathrm{X} ^{4}\Sigma^{+}$) and ground state atomic oxygen $(O(^3P))$.

Both ISC take place at the energies lower than that of the products making these exit channels practically barrierless in the reverse direction. Interestingly, the bent MSX11 structure occurs early on the decomposition pathway of i1 lying 123 kJ mol $^{-1}$ below the products, with the breaking Sn-O bond lengthened to 2.140 Å. Alternatively, the linear MSX12 structure is late, only 26 kj mol⁻¹ lower than the products and with breaking a Sn–O bond of 2.430 Å. The other exit MSX10 on the pathway from i4 to the SnO($X^1\Sigma^+$) + O(${}^{3}P$) products resides

82 kJ mol $^{-1}$ above the products making this channel noncompetitive. Singlet–triplet crossings also connect i1 with i2 (MSX1a (in SI) and MSX1b occurring in the close vicinity of i2), i1/i4 with i5 (MSX2 in the vicinity of singlet ts3), and i8 with i6 and i1 with i7 (MSX7 and MSX8, respectively, both located in the vicinity of singlet ts6). The other MSXs are responsible for internal conversion (IC) processes between different triplet PESs. For instance, MSX3 and MSX4 lead from i2 to i3 and i5, respectively, and reside lower in energy than i3 and i5. MSX5, MSX6, and MSX9 play the role of barriers for the interconversion of i3 to i5, i2 to i6, and i6 to i7, respectively.

Considering the energetics of the transition states, MSXs, and products, we can conclude that the most favorable reaction pathways include spin-allowed and adiabatic $Sn(^{3}P_j) + O₂(X³\Sigma_g⁻) \rightarrow$ ts1 \rightarrow i3 \rightarrow ts5 \rightarrow i7 \rightarrow SnO(X¹\Z⁺) + O(³P), spin-allowed nonadiabatic Sn(³P_j) + O₂(X ³ Σ_g^-) \rightarrow **i1** \rightarrow **MSX1b** \rightarrow **i2** \rightarrow **MSX3** \rightarrow **i3** \rightarrow ts5 \rightarrow i7 \rightarrow SnO(X¹\Z⁺) + O(³P), Sn(³P_j) + O₂(X³\Z_g⁻) \rightarrow ts2 \rightarrow i5 \rightarrow MSX5 $\rightarrow i3 \rightarrow t$ s5 $\rightarrow i7 \rightarrow SnO(X^{1}\Sigma^{+}) + O(^{3}P), Sn(^{3}P_{j}) + O_{2}(X^{3}\Sigma_{g}^{-}) \rightarrow$ $i1 \rightarrow ts6 \rightarrow MSS8 \rightarrow i7 \rightarrow SnO(X^{1}\Sigma^{+}) + O(^{3}P),$ and $Sn(^{3}P_{j}) +$ $O_2(X^3\Sigma_g^-) \rightarrow i1 \rightarrow ts6 \rightarrow MSSX7 \rightarrow i6 \rightarrow SnO(X^1\Sigma^+) + O(^3P),$ and spin-forbidden non-adiabatic $\text{Sn}(^{3}P_{j}) + \text{O}_{2}(X^{3}\Sigma_{g}^{-}) \rightarrow i1 \rightarrow$ MSX11 \rightarrow SnO(X¹ Σ ⁺) + O(³P) and Sn(³P_j) + O₂(X³ Σ_g^-) \rightarrow **i1** \rightarrow ts6 \rightarrow i8 \rightarrow MSX12 \rightarrow SnO(X¹ Σ ⁺) + O(³P). The competition between these channels can only be theoretically evaluated through cost-forbidden non-adiabatic *ab initio* molecular dynamics simulations involving multiple triplet and singlet PESs.

Conclusions

In conclusion, a merging of the electronic structure calculations and crossed beam experiments exposes the reaction dynamics in the tin $(Sn, {}^{3}P_{j})$ – molecular oxygen $(O_{2}, X^{3}\Sigma_{g}^{-})$ system forming tin monoxide (SnO, $X^1\Sigma^+$) along with atomic oxygen O(${}^{3}P$) in its electronic ground state. The reaction can be initiated on the singlet or triplet surfaces via additions of tin to the oxygen atom leading to linear, bent, or triangular structures. On both surfaces formation of a tin dioxide structure is required prior to unimolecular decomposition to SnO($X^1\Sigma^+$) and O(${}^{3}P$) on the triplet surface and SnO($X^1\Sigma^+$) with O(1D) on singlet PES. ISC and IC play important roles in this system and extensively interosculate the singlet and triplet surfaces via singlet–triplet (Fig. 5) and triplet–triplet (Fig. 5) intersections, respectively, via 13 minima on the seams of crossings (MSX). Thus, the starting channel on the triplet surface that leads to the barrier-free formation of triangular intermediate i2 $(SnO_2, C_{2v}, {}^{1}A_1)$ will be a cul-de-sac pathway in case of the absence of IC. What is interesting, a more thermodynamically competitive singlet surface without ISC channels has exit channels only to thermodynamically unfavorable $SnO(X^1\Sigma^+)$ and $O(^1D)$, which is not feasible under our collision energy of 13 kJ mol^{-1} . However, ISC offers direct barrier-free exit channels to $\text{SnO}(X^1\Sigma^+)$ and $O(^3P)$ products from i1 and i8 and the influence of the singlet surface on our experimental results cannot be excluded. The ISC is likely supported by the ''heavy atom effect'' of tin because the inclusion of heavy atoms in the molecular structure enhances the spin–orbit coupling between singlet and triplet states.^{74–77} In general, studied reactions of atomic oxidation for elements from XIV group ($E = Si$, Ge, Sn):^{49,50}

$$
E(^{3}P_{j})+O_{2}(X^{3}\Sigma_{g}^{-})\rightarrow \ EO(X^{1}\Sigma^{+})+O(^{3}P/^{1}D); \qquad (2)
$$

follows a similar mechanism, with a singlet surface having exit channels only to $EO(X^1\Sigma^+) + O(1D)$ and triplet to $EO(X^1\Sigma^+)$ + $O(^3P)$ without ISC. Within the Si \rightarrow Ge \rightarrow Sn series with increasing atomic radius: (i) enthalpy of reaction channels to $EO(X^1\Sigma^+) + O(1D)$ and to $EO(X^1\Sigma^+) + O(1D)$ increases with equal steps around \sim 140 kJ mol $^{-1}$ for each channel: -303 \rightarrow $-170 \rightarrow -24 \text{ kJ mol}^{-1} \text{ and } -107 \rightarrow +20 \rightarrow +166 \text{ kJ mol}^{-1},$ respectively; (ii) ISC effects enhances and expands.

Author contributions

Supervision and Funding acquisition – R. I. K., A. M. M; formal analysis – I. A. M.; investigation – I. A. M., Z. Y. and S. J. G. carried out the experimental measurements, A. A. N. – carried out the theoretical analysis; Writing original draft – I. A. M.; writing – review & editing – R. I. K., A. M. M., A. A. N., I. A. M.

Data availability

Essential data are provided in the main text and the ESI.† Raw data for crossed molecular beams experiment is available at https://doi.org/10.5281/zenodo.13831697.

Conflicts of interest

There are no conflicts to declare.

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