Enhanced electrical properties of antimony doped tin oxide thin films deposited via aerosol assisted chemical vapour deposition†

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Transparent conducting oxides have widespread application in modern society but there is a need to move away from the current ‘industry champion’ tin doped indium oxide (In₂O₃:Sn) due to high costs. Antimony doped tin(IV) oxide (ATO) is an excellent candidate but is limited by its opto-electrical properties. Here, we present a novel and scalable synthetic route to ATO thin films that shows excellent electrical properties. Resistivity measurements showed that at 4 at% doping the lowest value of 4.7 × 10⁻⁴ Ω cm was achieved primarily due to a high charge carrier density of 1.2 × 10²¹ cm⁻³. Further doping induced an increase in resistivity due to a decrease in both the carrier density and mobility. Ab initio hybrid density functional theory (DFT) calculations show the thermodynamic basis for the tail off of performance beyond a certain doping level, and the appearance of Sb(III) within the doped thin films.

Introduction

The commercial significance of producing efficient and sustainable transparent conducting oxides (TCOs) cannot be underestimated. The excellent optoelectronic properties of TCO thin films are essential for a number of applications including solar cells, gas sensors, catalysis, electronics and optics. 1,2 Thin films based on tin(IV) oxide have a wide fundamental band gap (~3.6 eV) and a stability comparable to other TCO systems making them a viable alternative to the more expensive indium-tin oxide (ITO) films.3 The enhancement of electrical n-type conductivity properties can be achieved by the introduction of intrinsic defects, including oxygen vacancies, or by introducing extrinsic defects through aliovalent dopants with, for example, higher valency cations.4 The electrical resistivity is dependent on the density of charge carriers and their mobility5 and for nominally undoped SnO₂ resistivity is limited to 10⁻²–10⁻³ Ω cm.1 Dopants such as fluorine and antimony have been used to great effect in improving the n-type conductivity of SnO₂ films,6–11 with the latter dopant producing blue coloured coatings. However, it is not desirable to dope beyond a certain concentration as this is known to increase electrical resistivity and reduce optical transparency.12,13 Three general considerations needed when doping are: (i) the solubility of the dopant in the structure; (ii) shallowness of the dopant level; (iii) the dopant should not behave as a compensatory acceptor (a low energy electron killing defect).14 Antimony-doped tin oxide (ATO) materials have received particular attention due to the availability of the constituent metals and a recent increase in the demand for architectured coloured glass.7 The production of doped tin oxide films has been achieved by a variety of physical and chemical vapour deposition methods.1,10 Aerosol-assisted chemical vapour deposition (AACVD) is a low-cost and versatile technique with the potential to be scaled up for the commercial production of thin films.15 Unlike, other methods, AACVD negates the need for highly thermally stable and volatile precursors which widens the choice available for producing films with tuneable properties.16

In this paper we doped tin(IV) oxide with varying concentrations of antimony. The as produced doped SnO₂ structure contains Sb in the +3 and +5 states with Sb(III) responsible for the bluish colouration of the films.6,8,17 Our ATO thin films show significantly enhanced electrical properties with resistivities as low as 4.7 × 10⁻⁴, and to our knowledge are on par or superior to other published studies, on this material. A computational analysis into the role of intrinsic and Sb-related defects in SnO₂ was carried out using hybrid density functional theory (DFT) and shows at low doping amounts, antimony is indeed incorporated into the lattice in the desired Sb(III) state.
Experimental

Deposition method

Depositions were carried in an atmosphere of air (BOC). The substrate used was SiO₂ barrier coated float glass (15 cm × 4 cm × 0.3 cm; Pilkington Ltd, UK). The coating (ca. 50 nm in thickness) prevents the ions within the glass diffusing to the surface. Any surface grease on the glass was removed using a washing regime of detergent, propan-2-ol and acetone. The precursor solution consisting of 0.30 g butyltin trichloride and dopant amounts of antimony(III) ethoxide (all from Sigma-Aldrich) in 10 mL methanol was placed in a glass bubbler. An aerosol was generated using a piezo electric device and carried, in a flow of air (1 L min⁻¹), through into horizontal bed cold-walled reactor (15 cm × 5 cm). The glass substrate, placed on the top and bottom of the reactor, was preheated to 450 °C on a graphite block containing a Whatman cartridge heater monitored by a Pt–Rh thermocouple. Depositions were achieved by the aerosol being carried in a stream of air through a brass baffle into the reactor. Waste vapours left through the exhaust. Once the entire precursor solution had been used up the reactor was allowed to cool to room temperature under air. Complete coverage of the substrate was achieved and the ATO films were blue and transparent with increasing dopant amounts in the precursor solution producing darker films.

Film analysis

X-ray diffraction (XRD) measurements were obtained using a modified Bruker-Axs D8 diffractometer with parallel beam optics fitted with a PSD LynxEye silicon strip detector to collect diffracted X-ray photons. X-rays were generated using a Cu source with Cu Kα1 and Cu Kα2 radiation of wavelengths 1.54056 and 1.54439 Å, respectively, with an intensity ratio of 2:1 and at 40 kV and 30 mA. The incident beam angle was kept at 1°, and the angular range of the patterns collected was 10° < 2θ < 66° with a step size of 0.05° counted at 4 s per step. The patterns were analysed for crystallinity and preferred orientation. Peak positions were compared to patterns from the Inorganic Crystal Structure Database (ICSD). The lattice parameters were calculated from powder X-ray diffraction data using the software GSAS and Expgui via the Le Bail method.

X-ray photoelectron spectroscopy (XPS) analysis of the films was carried out using a Thermo Scientific K-Alpha spectrometer equipped with a monochromatic Al-Kα source to identify chemical constituents. The peaks were modelled using CasaXPS software with binding energies adjusted to adventitious carbon (284.5 eV) in order to compensate for the effects of charging. Survey scans were collected in the range 0–1350 eV (binding energy) at a pass energy of 40 eV.

Wavelength dispersive X-ray (WDX) analysis was carried out on a Phillips ESEM. The Sn and Sb atomic% was derived from Sn-Kα line (25271.8 eV) and Sb-Kα (26359.5 eV), respectively.

UV/Vis/Near IR transmittance and reflectance spectra were produced using the Perkin Elmer Precisely Lambda 950 spectrometer using an air background and recorded between 320–2500 nm.

Scanning electron microscopy (SEM) was used to determine the film morphology from a top-down configuration using a JEOL JSM-6301F Field Emission instrument with accelerating voltages ranging from 3–5 keV on Au-coated samples. A side-on configuration was achieved using a JEOL JSM-6301F Field Emission instrument with accelerating voltages ranging from 5 keV on Au-coated samples.

Hall effect measurements were carried out on an Ecopia HMS-3000 set up in the Van der Pauw configuration to determine the sheet resistance, free carrier concentration (N) and mobility (μ). Samples of 1 cm² were prepared and silver paint (Agar Scientific) was used to form ohmic contacts which were tested on the in-built software prior to measurement. The samples were then subjected to an input current of 1 mA and a calibrated magnetic field of 0.58 T.

Theoretical methodology

A computational analysis into the role of intrinsic and Sb-related defects in SnO₂ was carried out using ab initio density functional theory (DFT) within the periodic code VASP. All geometrical and electronic optimisations were carried out using the hybrid functional PBE0 formalised by Adamo and Barone. PBE0 adds 25% of Hartree–Fock exchange to the PBE formalism resulting in an improved treatment of the self-interaction error compared to standard functionals. PBE0 can provide an accurate description of the band gap and geometry of SnO₂ compared to experiment, and has been shown to accurately describe the properties of other tin-based oxides. The Projector Augmented Wave method (PAW) was used to account for the interaction between the core electrons and the valence electrons of each species (Sn[Kr], O[He], Sb[Kr]). Initially the conventional unit cell of rutile SnO₂ was relaxed using a 400 eV plane-wave cutoff and a 4 × 4 × 6 Γ-centred k-point mesh to a force convergence tolerance of 0.01 eV atom⁻¹. A supercell expansion of 2 × 2 × 3 (72 atoms) was produced to probe the properties of each defect and its respective charge states. Structural optimisations were undertaken relaxing only the ions within the cell, whilst keeping the cell parameters constant utilising a Γ-centred k-point mesh of 2 × 2 × 2 wininned with a 400 eV plane-wave energy cut-off. All calculations were spin-polarised and convergence was deemed to be complete when the forces acting on all the ions was less than 0.01 eV atom⁻¹. Sb₂O₃, Sb₂O₄ and Sb₂O₅ were considered limiting phases in the formation of the dopant related species and were optimised using a plane-wave energy cut-off of 400 eV and Γ-centred k-point meshes of 2 × 5 × 4 (for the 28-atom Sb₂O₅ unit cell), 5 × 4 × 2 (for the 24 atom Sb₂O₄ unit cell) and 5 × 4 × 2 (for the 20 atom Sb₂O₃ unit cell). The volume, lattice parameters and atoms were allowed to relax until the forces acting on all ions was less than 0.01 eV atom⁻¹.

Defect formalism

The formation enthalpy for a defect in charge state ‘q’ was assessed using the relation:

\[
\Delta H_f(D, q) = (E^{0,q} - E^H) + \sum_i n_i (E_i + \mu_i) + q (E_{\text{Fermi}} + \epsilon_{\text{BM}} + \Delta E_{\text{pot}}) + q^2 E_{\text{corr}}^{1C} + E_{\text{corr}}^{BF}
\]
where $E^{D,q}$ is the total energy of the defective supercell in charge state $q'$ and $E^D$ is the total energy of the host supercell. $E_F$ and $\mu_i$ correspond to the elemental reference energies and the corresponding chemical potentials respectively (where $q'=Sn_{(s)},O_{2(g)}$ and $Sb_{(s)}$). $n$ is either positive or negative depending on whether the element is taken away from or added to the system respectively. $E_F$ is the Fermi energy and ranges from the valence band maximum (VBM) to $\sim 2.4$ eV above the conduction band minimum (CBM) which occurs at 3.6 eV. $\varepsilon_{\text{VBM}}$ is the eigenvalue of the VBM of the host material and is aligned to the defective supercell via a potential alignment term ($\Delta E_{\text{pot}}$) which averages the total potential of the supercell bar that in the immediate vicinity of the defect.\textsuperscript{35}

Due to the long ranged nature of the Coulomb interaction\textsuperscript{36,37} an image charge term $q^2e^{\text{IC}}$ was applied to account for the interaction of a charged defect and its periodic images. The image charge correction used in this work is based upon the Lany and Zunger formalism\textsuperscript{38} with the ‘non-cubic’ adaptation as implemented by Hine and Murphy.\textsuperscript{39} For shallow and resonant defects, a band-filling correction ($E_{\text{corr}}$) was applied to regain the dilute limit due to the relatively high carrier concentrations present in supercell calculations.\textsuperscript{38,40}

### Thermodynamic limits

A chemical potential range can be applied to account for the extremes in growth conditions; Sn-rich/O-poor and Sn-poor/O-rich. These are set within the bounds of the calculated formation enthalpy of SnO\textsubscript{2}:

$$\mu_{\text{Sn}} + 2\mu_{O} = \Delta H_{\text{f}}^{\text{SnO}_2} = -5.27 \text{ eV} \quad \text{(experiment) = -5.98 \text{ eV}}$$

(1)

A difference between calculation and experiment was expected due to the neglection of entropic effects within the athermal limit which the DFT calculations were performed. Thus the chemical potential limits for Sn and O ($\mu_{\text{Sn}}$ and $\mu_{O}$ respectively) for the two growth regimes were therefore:

Sn-rich/O-poor conditions: $\mu_{\text{Sn}} = 0 \text{ eV}; \mu_{O} = -2.64 \text{ eV}$  
Sn-poor/O-rich conditions: $\mu_{\text{Sn}} = -5.27 \text{ eV}; \mu_{O} = 0 \text{ eV}$

(2.1) and (2.2)

Further limitations were imposed on the system due to the formation of dopant related species. Fig. 1 displays the phase stability diagram within the constraints on $\mu_{O}$ (eqn (2.1) and (2.2)). Under Sn-rich/O-poor conditions ($\mu_{O} = -2.64 \text{ eV}$) the limitation on $\mu_{\text{Sn}}$ was based upon the formation of Sb\textsubscript{(s)} and therefore $\mu_{\text{Sn}} = 0 \text{ eV}$. Under Sn-poor/O-rich conditions ($\mu_{O} = 0 \text{ eV}$), $\mu_{\text{Sn}}$ was limited by the formation of Sb\textsubscript{2}O\textsubscript{5:41}

$$\mu_{\text{Sb}} + 2\mu_{O} = \Delta H_{\text{f}}^{\text{SbO}_5} = -9.61 \text{ eV} \quad \text{(experiment) = -9.97 \text{ eV}}$$

Therefore, $\mu_{\text{Sb}} = -4.80 \text{ eV}$ under Sn-poor/O-rich conditions.

The thermodynamic transition limits ($q/q'$) were defined as:

$$\varepsilon_{\text{D}}(q/q') = \frac{\Delta H_{\text{f}}(D, q) - \Delta H_{\text{f}}(D, q')}{q' - q}$$

The evolution of a defect from charge state $q$ to $q'$ at a certain Fermi level position can be seen experimentally in techniques such as deep level transient spectroscopy.

### Results and discussion

The films were deposited via AACVD using butyltin trichloride and antimony[III] ethoxide, when doped, in methanol at 450 °C using air as the carrier gas. The doped films were blue in coloration, which darkened with increasing dopant concentration. They had excellent coverage across the glass substrate, were adherent (confirmed by the Scotch\textsuperscript{TM} tape test) and air stable, showing no change over a one year period.

The successful incorporation of Sb into the SnO\textsubscript{2} films was confirmed by wavelength dispersive X-ray spectroscopy (WDX). The results show a positive correlation between the concentration of $[\text{Sb(OEt)}_3]$ in the precursor solution and the concentration in the films giving an average precursor efficiency of 47%.

X-ray diffraction (XRD) carried out on the pure and Sb doped SnO\textsubscript{2} films showed them to be phase pure and of the cassiterite phase with no peaks for SbO\textsubscript{x} observed even at high dopant concentrations therefore suggesting the successful formation of a solid solution (or a significant amorphous component). Le Bail modelling of the XRD patterns showed that doping of SnO\textsubscript{2} with Sb generally results in a contraction of the cassiterite unit cell (Table 1). This follows what is expected for the substitutional doping of Sb\textsuperscript{5+} as its ionic radii is only 60 pm whereas Sn\textsuperscript{4+} is 69 pm. At high Sb doping concentration (10 at%) there was an increase in unit cell size compared to 4 and 7 at%, which was most likely due to incorporation of Sb\textsuperscript{5+} (radii of 76 pm) ions as well as the desired Sb\textsuperscript{5+} into the lattice.

Upon doping a change in the preferred orientation was also seen with doped samples showing less preference for the (101) and (301) planes and more for (002) planes of cassiterite. Doping also caused a slight increase in the crystallite size as estimated using the Deby–Scherrer equation (Table 1). This is not normally observed with the presence of dopants, as the addition of foreign ions into the host lattice usually causes a distortion and hence a decrease in crystallinity. The general trend of increased crystallite size with doping concentration as

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**Fig. 1** Phase stability diagram for the Sb-based oxides under varying $\mu_{O}$ and $\mu_{\text{Sb}}$. $\mu_{O}$ ranges from 0 eV (the formation of O\textsubscript{2(g)}) to $-2.64$ eV (the formation of Sn\textsubscript{(s)}). Under O-rich conditions Sb\textsubscript{2}O\textsubscript{5} is the limiting phase and under O-poor conditions, Sn\textsubscript{(s)} is the limiting phase.
observed here maybe related to the increased film thickness (Fig. 2b–f). As these films were grown on amorphous glass substrates, layers in contact with and close to the substrate would be less crystalline and have smaller crystallite sizes. Due to templating effects the subsequent layers would then grow with larger crystallite sizes as these layers are now growing on crystalline layers beneath them. Therefore thicker films will on average have larger crystallite sizes. This effect has been previously observed for GaAs layers grown on glass via AACVD.42

Cross sectional high-resolution transmission electron spectroscopy (HRTEM) on the GaAs films found that crystallites close to the substrate were much smaller in size than the crystallites near the surface of the film.

All the films consist of overlapping tooth like features that are top heavy. It was generally found that thicker films had larger widths of features at the top. Apart from this, the morphology of the films remained fairly consistent with thickness. Notably only one type of crystal growth was seen with no evidence of any amorphous features, strengthening the argument for a complete solid solution.

X-ray photoelectron spectra (Fig. 3a) showed the Sn 3d5/2 peak to correspond to literature examples of Sn(IV) and to be

<table>
<thead>
<tr>
<th>Film</th>
<th>Film thickness (nm)</th>
<th>Crystallite size (nm)</th>
<th>a (Å)</th>
<th>c (Å)</th>
<th>Unit cell volume (Å3)</th>
<th>χ²</th>
<th>Fitted wR_p</th>
</tr>
</thead>
<tbody>
<tr>
<td>SnO2</td>
<td>480</td>
<td>16</td>
<td>4.7670(5)</td>
<td>3.1902(2)</td>
<td>72.493(14)</td>
<td>2.1</td>
<td>17</td>
</tr>
<tr>
<td>Sn0.99Sb0.01O2</td>
<td>550</td>
<td>23</td>
<td>4.7687(6)</td>
<td>3.1945(2)</td>
<td>72.643(23)</td>
<td>2.4</td>
<td>18</td>
</tr>
<tr>
<td>Sn0.96Sb0.04O2</td>
<td>525</td>
<td>19</td>
<td>4.7510(4)</td>
<td>3.1935(2)</td>
<td>72.081(11)</td>
<td>1.7</td>
<td>14</td>
</tr>
<tr>
<td>Sn0.93Sb0.07O2</td>
<td>800</td>
<td>25</td>
<td>4.7510(4)</td>
<td>3.1915(2)</td>
<td>72.041(11)</td>
<td>2.0</td>
<td>17</td>
</tr>
<tr>
<td>Sn0.90Sb0.10O2</td>
<td>800</td>
<td>23</td>
<td>4.7614(3)</td>
<td>3.1944(1)</td>
<td>72.421(8)</td>
<td>2.2</td>
<td>16</td>
</tr>
</tbody>
</table>

Fig. 2 (a) XRD patterns of the as deposited and nominally Sb doped SnO2 thin films grown via AACVD. (b–f) Show the cross-sectional SEM images with the top down images as inset (scale bar for all images is equal to 500 nm).

Fig. 3 (a) Core level XPS spectra for the Sn 3d and Sb 3d signals. (b) Valence band spectra for all the AACVD grown films.
centred at 486.5 eV for all films apart from Sn$_{0.96}$Sb$_{0.04}$O$_2$ that showed the peak to be slightly shifted to 486.1 eV.

The 3d$_{3/2}$ peak for Sb overlaps with the O 1s but the less intense 3d$_{5/2}$ can be used to confirm the presence of Sb in the films. The overlapping Sb 3d$_{3/2}$ and O 1s peak was modelled with a spin orbit split of 9.39 eV and the Sb 3d$_{5/2}$ peak intensity was 1.5 times greater than the Sb 3d$_{3/2}$. Due to the similar peak positions of Sb(V) and Sb(III) it is difficult to differentiate between these two oxidation states with certainty from XPS alone, especially at dopant concentrations where signal to noise in the spectra can be low.

Valence band (VB) photoelectron spectra for the pure and Sb doped SnO$_2$ films are shown in Fig. 3b. All spectra show the three peaks made up of O 2p states that are expected for the VB structure of SnO$_2$. The overall O 2p band width is roughly 9 eV. Apart from some peak broadening there was not much change in the VB due to Sb doping. No electronic states were detected within the band gap as a result of doping, this is unusual as normally a peak is expected 0.5 eV below the Fermi level indicating conduction band filling upon semiconduction to metal transition. The absence of this peak maybe due to Sb being primarily a bulk dopant in the AACVD grown films. This also follows from core level XPS and WDX results that showed Sb levels at the surface being much lower than in the bulk therefore suggesting bulk segregation.

Hall effect measurements were taken at room temperature using the van der Pauw technique (Table 2). All the films showed n-type conductivity from the negative Hall coefficients. The resistivity, mobility and bulk concentration (charge carrier density) of undoped SnO$_2$ were $1.8 \times 10^{-3}$ Ω cm, 10.4 cm$^2$ V$^{-1}$ s$^{-1}$ and $3.4 \times 10^{20}$ cm$^{-3}$ respectively, values which in general are in the ranges quoted in the literature, albeit, having produced the films through a range of techniques.$^{1,4,44}$

The n-type conductivity is attributed to the formation of shallow donor levels occurring due to the presence of oxygen vacancies and interstitial tin in the SnO$_2$ lattice.$^{4,44}$ While the resistivity increased when 1 at% Sb was present in the SnO$_2$ film, a doping level of 4 at% reduced the resistivity by a factor of 10. This was associated with an increase in mobility and carrier concentration which can be attributed to the ratio of Sb$^{5+}$ and Sb$^{3+}$ present in the lattice.$^{46}$ The Sb$^{5+}$ state contributes additional free electrons into the conduction band increasing conductivity and the charge carrier concentration.$^{47}$ Above 4 at% Sb the resistivity increased with a corresponding decrease in mobility and carrier concentration. In such cases, more of the Sb$^{5+}$ ions reduce to the Sb$^{3+}$ state which act as acceptors hence removing electrons from the conduction band.

Scattering of charge carriers limits mobility in polycrystalline materials.$^{47,48}$ A possible reason for the decrease in carrier concentration in excess of 4 at% is due to the increased disorder that results from the donor having an increased activation energy.$^{47}$

The transmittance and reflectance spectra of the films were collected using visible/near IR spectroscopy between 320 and 2500 nm and are shown in Fig. 4. The undoped SnO$_2$ film had the highest transmittance of up to 85% in the visible region. Doping Sb reduced the optimum transmittance in the visible range and increased reflectance in the infra-red. Transmittance decreased as a function of dopant level (from a maximum of 80% down to 45%). Correspondingly, the intensity of the blue colouration of the films increased which is commonly reported.$^{5,17,49}$ However, a variety of explanations have been put forward to account for this observation to fit in with the opto-electronic properties of the films reported. One suggestion is that increasing the Sb dopant amounts leads to an increase

![Fig. 4 The transmittance (black) and reflectance (red) spectra of films produced via AACVD.](image)

### Table 2 The electrical properties of the pure and Sb doped films as determined via room temperature Hall effect measurements

<table>
<thead>
<tr>
<th>Film</th>
<th>Film thickness (nm)</th>
<th>Sheet resistance (Ω sq$^{-1}$)</th>
<th>Resistivity, $\rho$ (Ω cm)</th>
<th>Mobility, $\mu$ (cm$^2$ V$^{-1}$ s$^{-1}$)</th>
<th>Charge carrier concentration, $N$ (cm$^{-3}$)</th>
</tr>
</thead>
<tbody>
<tr>
<td>SnO$_2$</td>
<td>480</td>
<td>37.1</td>
<td>$1.8 \times 10^{-3}$</td>
<td>10.4</td>
<td>$3.4 \times 10^{20}$</td>
</tr>
<tr>
<td>Sn$<em>{0.99}$Sb$</em>{0.01}$O$_2$</td>
<td>550</td>
<td>46.1</td>
<td>$2.5 \times 10^{-3}$</td>
<td>6.9</td>
<td>$3.6 \times 10^{20}$</td>
</tr>
<tr>
<td>Sn$<em>{0.96}$Sb$</em>{0.04}$O$_2$</td>
<td>525</td>
<td>9.0</td>
<td>$4.7 \times 10^{-4}$</td>
<td>11.4</td>
<td>$1.2 \times 10^{21}$</td>
</tr>
<tr>
<td>Sn$<em>{0.93}$Sb$</em>{0.07}$O$_2$</td>
<td>800</td>
<td>20.2</td>
<td>$1.6 \times 10^{-3}$</td>
<td>6.2</td>
<td>$6.2 \times 10^{20}$</td>
</tr>
<tr>
<td>Sn$<em>{0.90}$Sb$</em>{0.10}$O$_2$</td>
<td>880</td>
<td>39.1</td>
<td>$3.4 \times 10^{-3}$</td>
<td>3.1</td>
<td>$5.8 \times 10^{20}$</td>
</tr>
</tbody>
</table>
in excitable electrons attributed to a greater presence of Sb$^{5+}$.\textsuperscript{49} In contrast, others suggest, based on the observed reduction in conductivity that the darkening must be due to the presence of an increase in Sb$^{3+}$ ions concentration.\textsuperscript{50}

The plasma edge was estimated from the energy at which the reflectivity reaches half its maximum value.\textsuperscript{3} In general, the plasma band onset of increased reflection shifted towards shorter wavelengths for the doped films compared with pure SnO$_2$.

Emissivity is the ability of a material to emit infrared radiation; values range from 0 for a shiny mirror to 1 for a matt black body. For heat mirror applications reflections in the visible and near infrared are important. A low emissivity value is ideal for a heat mirror as reflection is high in the near infrared. The best commercially available TCO having heat mirror properties is Pilkington’s fluorine tin oxide (FTO) having emissivity value of 0.15. Emissivity normally drops with increasing when more dopant is added. The emissivity of the undoped SnO$_2$ film was 0.33. Doping with 1 at% Sb reduced the emissivity to 0.23, reaching a minimum of 0.22 at a dopant concentration of 7 at% and then increasing to 0.30 at the highest dopant concentration.

### Computational analysis

\textit{Ab initio} density functional theory (DFT) was used to determine the role of both intrinsic and Sb dopant related defects on the electrical properties of SnO$_2$.

The thermodynamic transition levels are displayed in Fig. 5 and show the growth conditions under Sn-rich/O-poor (typically n-type favourable conditions) and Sn-poor/O-rich (p-type preferred conditions) and represent the extreme bounds of the equilibrium growth conditions. The dominant acceptor and donor intrinsic defects were modelled to aid the understanding of the Sn-related defects.

The oxygen and tin vacancies (V$_O$ and V$_Sn$) represent the dominant intrinsic donor and acceptor defects present in SnO$_2$.\textsuperscript{54} V$_0$ is a low formation energy deep donor under Sn-rich/O-poor conditions, where the formation energy of the neutral charge state is $\sim 1.89$ eV and rises to $\sim 4.53$ eV under Sn-poor/O-rich conditions.

As in other n-type oxides, such as ZnO$^{31,51-53}$ and BaSnO$_3$,\textsuperscript{29} V$_O$ is a negative-U defect with the 2+/0 transition level occurring $\sim 0.76$ eV below the CBM in SnO$_2$ revealing that V$_O$ will not contribute to the conductivity, which is supported by theoretical\textsuperscript{54,55} and experimental\textsuperscript{56} results alike. V$_Sn$ also acts as a very deep acceptor with the 0/1$^-$ transition level occurring $\sim 1.75$ eV above the VBM. V$_Sn$ has a very high formation energy over the range of the band gap with the formation energy under the most favourable growth conditions (Sn-poor) being prohibitively high at $\sim 8.37$ eV.

Sb$_{Sn}$ is a resonant donor with the 1+/0 transition level occurring $\sim 0.15$ eV above the CBM and has a low formation energy under both Sn-rich/O-poor and Sn-poor/O-rich growth regimes. Interestingly, Sb$_{Sn}$ is highest in energy ($\Delta H_{[Sb_{Sn}^0]} = \sim 1.42$ eV) under typically n-type favourable “O-poor” conditions and lowest under a p-type favourable “O-rich” regime ($\Delta H_{[Sb_{Sn}]} = \sim 0.95$ eV). This is due to the lower $\mu_{Sn}$ under Sn-poor/O-rich conditions and the availability of free Sn sites. The facile incorporation of Sb, as found with the AACVD films here, is likely due to the similar ionic radii to Sn ($Sn = 0.69$ Å; $Sb = 0.6(Sb^{5+})–0.76(Sb^{3+})$ Å)\textsuperscript{61} and over all charge states Sb is shown to have little distortion on the rutile lattice. Fig. 6a shows the partial charge density for Sb$_0^0$ displaying a localization of the electron density in the Sb 5s orbital to the Sb defect. Under both growth conditions, the 1−/− transition level occurs $\sim 0.41$ eV above the CBM and involves an extra electron being trapped on the Sb 5s orbital forming Sb$^{5+}$\textsuperscript{60} (Fig. 6b). As the defect states occur at the conduction band minimum there is small localization of density on O s states and Sn s states. Previous hybrid DFT calculations support a resonant low formation energy donor under both conditions,\textsuperscript{62–64} however Lany and co-workers propose that Sb is a negative U defect existing in only the 1− and 1+ charge states in contrast to our work.\textsuperscript{64}

The Sb–O bond lengths increase from $\sim 2.0$ Å to $\sim 2.03$ Å from the neutral to the 1− charge state so little distortion is seen. Under Sn-poor/O-rich conditions, Sb$_0^0$ is compensated for by V$_{Sn}^-$ $\sim 0.45$ eV above the CBM and at $\sim 2.05$ eV under Sn-rich/O-poor conditions. Under both growth extremes therefore, the conductivity and mobilities in ATO films will be limited by the formation of the 0 and 1− charge state.

Extended X-ray absorption fine structure (EXAFS) have proposed the clustering of Sb$_{Sn}$ and V$_O$ leading to the formation of Sb$^{11}$ and a trapped electron in the V$_O$ site.\textsuperscript{65,66} [Sb$_{Sn}$ + V$_O$] clusters were produced for this study in both a ‘near’ and ‘far’ configuration to examine this effect. The near configuration involved Sb$_{Sn}$, neighbouring a V$_O$ whilst the far configuration separated the two defects by $\sim 8$ Å. From our calculations, we find that Sb$_{Sn}$ prefers to distort from its Sn site towards the oxygen vacancy with a binding energy of $\sim 0.78$ eV. As such, higher Sb doping concentrations will facilitate the association of Sb with an oxygen vacancy. Typically, Coulombic effects should deter this behaviour from occurring considering that n-type defects should repel each other, however, this clustering is stabilised by the electronic rearrangement to give a lone pair on the Sb site. [Sb$_{Sn}$ + V$_O$] in the far configuration is less energetically favourable than [Sb$_{Sn}$ + V$_O$] in the near configuration by $\sim 0.4$ eV.
Under Sn-rich/O-poor conditions, \([\text{Sb} + \text{V} + \text{O}]\) is a shallow donor with a formation energy of \(\sim 3.03\) eV, this rises to \(\sim 5.20\) eV under an Sn-poor/O-rich regime. \([\text{Sb} + \text{V} + \text{O}]\) acts as a one electron donor and has two transition levels deep in the bandgap, the 3+/2+ and 2+/1+ transition levels occur \(\sim 1.75\) eV and \(\sim 1.59\) eV below the CBM respectively. The 1+/0 transition level occurs \(\sim 0.48\) eV above the CBM. The partial charge densities for the neutral, 1+ and 2+ charge states are shown in Fig. 7a, b and c respectively. For the neutral charge state \([\text{Sb} + \text{V} + \text{O}]^0\), two electrons are located on a lone pair on Sb (indicating Sb(III)) with the remaining electron in the VO. This is consistent with the EXAFS analysis in which 5-coordinate Sb is seen corresponding to a removal of a neighbouring oxygen.\(^{65,66}\) In the 1+ charge state an electron is removed from the vacancy site leaving the two electrons in the Sb lone pair. The 2+ and 3+ charge states involve the subsequent removal of electrons from the lone pair as shown in Fig. 7b and c. As the 3+ and 2+ charge states are very deep in the band gap it is unlikely that these will be seen and as such there will be minimal effect on the optical transparency. It is therefore likely that Sb(III) will be seen for most situations where the Fermi level is greater than 2.01 eV. Sb hardly moves from its position over the range of charge states (maximum distortion of 2% in all three directions) despite the Sb ion originally distorting slightly towards the oxygen vacancy.

Under both growth regimes, Sb is a very high formation energy donor, and is unlikely to form at Fermi levels near the CBM. Under Sn-rich/O-poor and Sn-poor/O-rich conditions, the formation energy of Sb\(^+\) is \(\sim 9.40\) eV and \(\sim 14.20\) eV respectively, and so quantities of Sb are expected to be negligible.

Sb is therefore a low formation energy donor under both doping regimes and is expected to incorporate as Sb(III) at low doping concentrations. Towards higher concentrations, Sb may either form a localised polaron or likely have a preference to associate with an oxygen vacancy (VO) forming Sb(III) provided...
the concentration of oxygen vacancies are substantial. This defect complex, however, still donates an electron to the conduction band, accounting for the high conductivities in Sb-doped SnO$_2$ samples here and elsewhere$^{67}$ as well as the prevalence of Sb(III) and Sb(V) in the SnO$_2$ lattice.

**Conclusion**

The AACVD prepared cassiterite phase pure ATO thin films using simple precursors at 450 °C show excellent optoelectrical properties. The best performing sample (4 at% Sb doped) showed an electrical resistivity of $4.7 \times 10^{-4}$ $\Omega$ cm resulting from an electron mobility of 11.4 $\text{cm}^2\text{V}^{-1}\text{s}^{-1}$ and a relatively high carrier density of $1.2 \times 10^{21}$ $\text{cm}^{-3}$. These results are comparable to some of the lowest resistivity values to date presented in literature for ATO hence having promising technological applications in optoelectronic devices. DFT calculations showed that at low doping concentrations, Sb incorporates into the SnO$_2$ lattice as Sb(v). Sb(v) is a shallow donor with low formation energy. At higher doping regimes, the calculations show that substitutional Sb reduces to Sb(III) or associates with oxygen vacancies producing Sb(III) [present in the AACVD films] acting as a potential limitation to ATO thin films.

**Conflicts of interest**

There are no conflicts to declare.

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**References**


