Solution phase growth and analysis of super-thin zigzag tin selenide nanoribbons

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Abstract
Tin selenide (SnSe), a highly promising layered material, has been garnering particular interest in recent times due to its significant promise for future energy devices. Herein we report a simple solution-phase approach for growing highly crystalline layered SnSe nanoribbons. Polyvinylpyrrolidone (PVP) was used as a templating agent to selectively passivates the (100) and (001) facets of the SnSe nanoribbons resulting in the unique growth of nanoribbons along their b-axis with a defined zigzag edge state along the sidewalls. The SnSe nanoribbons are few layers thick (∼20 layers), with mean widths of ∼40 nm, and achievable length of >1 μm. Nanoribbons could be produced in relatively high quantities (>150 mg) in a single batch experiment. The PVP coating also offers some resistance to oxidation, with the removal of the PVP seen to lead to the formation of a SnSe/SnOₓ core-shell structure. The use of non-toxic PVP to replace toxic amines that are typically employed for other 1D forms of SnSe is a significant advantage for sustainable and environmentally friendly applications. Heat transport properties of the SnSe nanoribbons, derived from power-dependent Raman spectroscopy, demonstrate the potential of SnSe nanoribbons as thermoelectric material.

Supplementary material for this article is available online

Keywords: nanoribbon, tin selenide, Raman spectroscopy, 1d van-der Waals structure

(Some figures may appear in colour only in the online journal)

Introduction
Quasi-one-dimensional (1D) nanoribbon structures, formed from two-dimensional (2D) layered materials, have exciting and unique physical properties compared to their conventional 2D, or bulk counterparts. Engineering nanostructures into these confined nanoribbon morphologies, where length > width > -thickness, is especially relevant for layered materials due to the added advantage of shape and size-dependent optical and...
of SnSe, which is not attainable with other morphological features. To this end, new and easy to implement synthetic methods for the growth and morphological engineering of SnSe nanoribbons are required.

Many solution-based methods exist for the synthesis of metal chalcogenide materials, with advantages lying in the ease and scalability of these methods versus costly vapour deposition based methods. Though other SnSe nanostructures such as nanoparticles, [28] nanorods, [29] large nanobelts [30] and 2D like flakes [31, 32] have previously been synthesised in solution using directing and capping agents such as oleylamine, ethanamine and 1,10-phenanthroline, growth of SnSe nanoribbons is still elusive in any solution phase processing. Additionally, developing a green synthesis method for the high yield and controllable synthesis of these materials is of paramount importance if they are to be successfully incorporated into future electronic and energy devices. Polyvinylpyrrolidone (PVP) is an alternative nontoxic and bio-compatible surface capping and structure-directing agent, that is commonly used for the controlled synthesis of nanoparticles [33] and nanowires, [34] due to its non-toxic nature [35]. PVP is known to offer selective shape control in solution grown crystalline materials, offering the tailored growth of nanowire structures by selective binding to different crystal planes [36]. To the best of our knowledge, PVP has never been used as a templating agent to grow SnSe or other group IV monochalcogenides in 1D nanoform. PVP has been shown to demonstrate biocompatibility in combination with SnSe, [37] where it was used to surface passivate SnSe nanorods in a post-growth treatment. However, passivation of SnSe with PVP during growth was not studied previously.

While SnSe nanowires have been previously explored, through both chemical vapour deposition [12, 38, 39] and solution based methods [40, 41], herein we report for the first time, the simple, environment-friendly and scalable solution phase synthesis of super-thin (∼20 layers thick) SnSe nanoribbons. The nanoribbons are single crystalline throughout and are synthesised using a simple and scalable approach with readily available precursor compounds and PVP as a templating agent. SnSe nanoribbons were structurally analysed in detail via high-resolution transmission electron microscopy. SnSe nanoribbons have a zigzag structure with a (010) growth direction (along b-axis), which is highly relevant for thermoelectric applications. A laser-induced heat generation and heat transport study for nanoribbons during Raman measurements, by analysing the Raman peak position, portray SnSe nanoribbons as a promising thermoelectric material.

**Experimental**

SeO₂ (99.8%), polyvinylpyrrolidone (PVP, average molecular weight = 40 000 kDa), KOH, ethylene glycol (emplura), SnCl₂:2H₂O (≥ 99.99%) and ascorbic acid (≥ 99%) were all purchased from Sigma Aldrich.
**Synthesis of (Se)\textsubscript{6}(PVP)\textsubscript{8} nanoparticles**

SnSe nanoribbons were synthesised by first preparing a solution of (Se)\textsubscript{6}(PVP)\textsubscript{8} nanoparticle complexes. In a typical synthesis, 0.3327 g of SeO\textsubscript{2} (3 mmol), 3 g PVP and 0.675 g KOH were dissolved in 45 ml of ethylene glycol by heating to 80 °C under Ar while stirring, yielding a very light yellow solution. After complete dissolution the temperature was increased to 195 °C for 18 h, giving a black brown-reddish solution.

**Synthesis of SnSe nanoribbons**

To synthesise SnSe nanoribbons, two additional solutions were then prepared: (1) 3 mmol SnCl\textsubscript{2}.2H\textsubscript{2}O (0.675 g) and 0.3 g PVP were dissolved in 12 ml ethylene glycol at 80 °C under stirring and (2) 6 mmol of ascorbic acid (1.0566 g) in 4 ml ethylene glycol was prepared by heating to 80 °C under stirring. These two solutions were then mixed and injected together into the previously prepared solution of Se-PVP nanoparticles in ethylene glycol at 80 °C under stirring. Upon addition of the SnCl\textsubscript{2}-PVP-ascorbic acid mixture the solution undergoes a colour change to deep red-brownish; stirring was maintained for 5 min at 80 °C. This solution was then added to a Teflon lined autoclave (~7.5 ml per 20 ml cell volume) and heated at 195 °C for between 27 and 76 h. After cooling to room temperature, the solution was washed twice with deionised water followed by centrifugation and then re-dispersed in ethanol.

**Characterisation**

The synthesised nanomaterials were analysed using a Helios Nanolab 600i scanning electron microscope (SEM). Raman spectroscopy data was taken with a Horiba Jobin Yvon micro-Raman spectrometer, which is equipped with 1800 lines/mm grating and a peltier cooled CCD. An Ar\textsuperscript{+} laser of wavelength 488 nm was used to excite the sample and a 50× objective was used to focus the laser on sample as well as to collect the data in backscattering geometry. The crystal structure of the product was confirmed by x-ray diffraction (XRD) using a Philips X’pert Pro MPD equipped with a Panalytical Empyrean Cu x-ray tube and a Philips X’celerator detector. X-ray photoelectron spectroscopy (XPS) analysis was undertaken on an Oxford Applied Research Escabase XPS system with a base pressure of 5 × 10\textsuperscript{–10} mbar and a non-monochromated Al Kα x-ray source at 200 W. Survey spectra were recorded between 0 and 1000 eV at a step size of 0.7 eV, a dwell time of 0.3 s and a pass energy of 100 eV. Core levels scans were acquired at a step size of 0.1 eV, a dwell time of 0.1 s and a pass energy of 20 eV. CasaXPS software was used for spectra processing with peaks corrected to a Shirley background and fitted to Voigt profiles. Charge correction was applied to the C 1s peak at 285 eV. Samples were dropcast onto a carbon coated TEM grid for detailed microscopy analysis. TEM analysis was performed on a JEOL 2100 electron microscope operating at 200 kV. EDX elemental mapping was performed at 30 kV on a Helios Nanolab 600i. High annular dark field and bright field STEM imaging and electron energy loss spectroscopy (EELS) was performed on a Probe Corrected JEOL ARM TEM operating at 200 kV. Vesta software was used for the illustration of crystal structures [42].

**Results and discussion**

SnSe nanoribbons were synthesised through a PVP-mediated solution phase method in a Teflon-lined autoclave (see the experimental section for a detailed procedure). Figure 1 highlights how the morphology of the nanoribbons varies as a function of reaction time at a growth temperature of 195 °C. The product obtained after 27 h of reaction resembles short 1D nanostructures as seen in figure 1(a). EDX analysis revealed these to consist of approximately ~1:1 Sn:Se (see supplementary information figure S2). The mean lengths and widths of these structures were found to be 216 ± 119 nm and 37 ± 13 nm respectively. A negligible amount of 2D flakes were seen in this reaction. Increasing the reaction time to 50 h resulted in an increase in the mean lengths of the 1D structures but not their widths, i.e. a mean length of 358 ± 185 nm and width of 36 ± 17 nm was obtained. After a 27 h reaction time, 90% of the nanoribbons had lengths < 400 nm, but after 50 h, 62% of the nanoribbons had lengths < 400 nm, with 90% of them below 650 nm; thus indicating that the mean length of the nanoribbons is directly proportional to the reaction time (further statistical analysis is detailed in supplementary information figure S3). Increasing the reaction time from 27 to 50 h therefore resulted in an increase of the mean nanoribbon length by 189 nm. However, the nanoribbon widths remained more or less constant with the increase in the reaction time, with only a negligible increase of ∼1 nm in the mean width being observed for 27 h compared to 50 h reaction. Of note, interestingly, the reactant solution was seen to contain small nanoparticles before transferring to the autoclave for nanoribbon growth (see supplementary information figure S4 for SEM of the dropcasted Se-PVP solution, and figure S5 for the SEM of the dropcasted Se-PVP + SnCl\textsubscript{2}-PVP mixture). The increase in reaction time thus develops the nanostructures from the initial nanoparticles to 1D nanoribbons, which further increase in length with increasing reaction time. The longest nanoribbons still maintained a mean width similar to the shorter nanoribbons, thus allowing the facile tuning of the aspect ratio with reaction time.

Facilitating the future implementation of SnSe nanoribbons into devices, such as Li-ion batteries and thermoelectric generators, requires production of the nanomaterial in reasonable quantities. Scaling up of 1D SnSe nanorod reactions while...
maintaining a length:width aspect ratio >5 has previously proved difficult [43]. The batch synthesis of SnSe nanoribbons produced in our approach was found readily scalable, capable of producing hundreds of mg of 1D SnSe nanoribbons with length:width aspect ratios of up to ∼40 achievable (see supplementary information figure S6). To demonstrate the scalability of the growth process, the reaction volume was increased while keeping the reaction temperature and time constant. Figure 2(a) shows the morphology of the structures formed when the reaction volume was roughly doubled to 45 ml vessel, with the reaction temperature kept at 195 °C and the reaction time set at ∼3 d (76 h). The mean length and width of these nanoribbon structures were observed to be 439 ± 211 nm and 42 ± 10 nm respectively, which were consistent with the increase in length with reaction time trend seen for the nanoribbons obtained in smaller 22 ml reaction vessels (figure 1). The nanostructures obtained upon scaling up to a 125 ml reaction vessel at 195 °C for 3 d (72 h) are shown in figure 2(c).

A large amount of SnSe nanostructures (∼158.7 mg) was obtained from the reaction in the 125 ml vessel. At this higher reaction volume a small increase in the ratio of 2D flakes to 1D nanoribbons was observed, giving an 84% yield of nanoribbons to flakes in the 125 ml reaction vessel (see supplementary information figure S6), suggesting that adjustment of the reaction conditions (such as PVP to reactant ratio) may be necessary for higher reaction volumes. Figure 2(d) shows the statistical analysis of the nanostructures obtained in the high volume reaction, revealing a larger spread in the lengths and widths (554 ± 258 nm mean length and 118 ± 129 nm mean width) due to the 2D flakes present in the sample. A small increase in nanostructure length is also observed at the higher reaction volume. After excluding the 2D flakes while analysing the width of the nanoribbon structures, mean width of 51 ± 18 nm was observed for the nanoribbons, relatively similar to that achieved for lower reaction volumes. The aspect ratio of the produced 1D nanoribbons was greatly enhanced
through our PVP-assisted approach compared to previously reported solution-grown SnSe nanorods.

Figure 3(a) shows a typical XRD pattern for the SnSe nanoribbons. Phase pure orthorhombic Pnma 62 SnSe was seen, with the 200 and 400 peaks highly prominent. No indication of the presence of any other SnSex phases such as SnSe2 was seen in the XRD, proving the high phase purity of the SnSe nanoribbons synthesised. Figure 3(b) shows a Raman spectrum taken from a sample of the SnSe nanoribbons (after 76 h in the 45 ml vessel). Clear peaks in the spectrum are seen at 77, 105, 127 and 157 cm\(^{-1}\), which can be assigned to the A\(_{1g}\), B\(_{3g}\), A\(_{2g}\) and A\(_{3g}\) Raman modes of SnSe respectively. At room temperature, bulk SnSe belongs to the Pnma (No. 62) space group (orthorhombic Bravais lattice) and it is expected for Pnma SnSe to possess 12 Raman active modes \cite{44}. Experimental Raman spectra recorded from SnSe nanoribbons depicted four Raman modes (3 A\(_g\) and 1 B\(_{3g}\)\(_g\)) similar to the ones observed for ultrathin SnSe flakes, SnSe nanolayers and single crystalline SnSe nanosheets \cite{45–47}. However, compared to previously reported Raman analysis for SnSe flakes and 2D layers, different shifts in the Raman modes has been observed. For example, no shift in the B\(_{3g}\) Raman mode was observed in comparison with the 25 nm thick SnSe nanosheets, \cite{47} whereas a 3 cm\(^{-1}\) red shift and 2 cm\(^{-1}\) blue shift in the B\(_{3g}\) Raman mode was observed in the SnSe nanoribbon sample compared to previous nanoflake and 2D layered (21 layers) structures, respectively \cite{45, 46}. If we compare the Raman modes from the nanoribbons with the theoretically predicted Raman modes for bulk SnSe, the A\(_{1g}\) and A\(_{3g}\) modes are blue shifted by 6 cm\(^{-1}\) whereas the A\(_{2g}\) and B\(_{3g}\) modes are red-shifted by 3 cm\(^{-1}\) \cite{44, 48}. The observed Raman shift relative to the bulk and other 2D nanostructures could potentially result from the morphology, shape asymmetry, nanostructure thickness, and anisotropic crystal confinement along the c-axis for the SnSe nanoribbons.

Nanoribbons made of few-atom thick lamellar crystals are novel forms of 1D nanomaterials and are ideal systems for investigating the size and dimensionality dependence of fundamental properties. We have also performed a laser power dependent Raman study on a SnSe nanoribbon sample by varying the laser (Ar\(^{+}\) 488 nm) power from 0.01 to 0.08 mW to explore the thermophysical properties of the nanoribbon structure. The corresponding Raman spectra of the SnSe nanoribbons with varying laser power are displayed in figure 3(c). To get the peak positions and full width at half maxima, the spectra were fitted with Lorentzian function. It is very clear that the peak positions and peak intensity of all the Raman modes for SnSe nanoribbons change with laser power. The increase in the Raman intensity is due to the increase in photon flux with the laser power. Figure 3(d) shows the variation of peak positions as a function of laser power for the different Raman modes of SnSe nanoribbons. For reference, the same measurement was performed on bulk Si (shown in figure S7 in the supporting information). The peak positions for all the modes associated with SnSe decrease monotonically with increasing laser power, which indicates an
increase in the local temperature of the sample due to laser irradiation \([49, 50]\). The factors that influence the temperature dependence of Raman shifts and line widths are anharmonic phonon–phonon interaction and thermal expansion. To analyse the red-shift of the Raman modes with laser power we used a linear change in energy: \(\omega_P = \omega_0 + \chi P\), where \(P\) is the laser power, \(\omega_P\) is the peak position at laser power, \(P\) and 
\[\chi = \frac{\partial \omega}{\partial P}\] is the measure of the rate of change of peak position with laser power. By fitting the phonon downshift using this equation, the obtained \(\chi\) values for different SnSe Raman modes vary from 17.5 to 96.4 \(\text{cm}^{-1} \text{mW}^{-1}\) (see table 1 in the supporting information), whereas for the bulk Si \(\chi\) value is found to be 0.96 \(\text{cm}^{-1} \text{mW}^{-1}\), which is in agreement with the previous reports \([51]\). The large red shifts in the SnSe Raman modes with increasing laser power (compared to bulk Si) indicates a high rise in the local temperature for SnSe nanoribbons compared to bulk Si. The high \(\chi\) value in SnSe compared to Si signifies the slow heat transfer (as thermal conductivity \((\kappa)\) is inversely proportional to \(\chi\) \([52]\)) within the SnSe nanoribbon sample because of the low thermal conductivity of the nanoribbons. These results demonstrate the

**Figure 3.** (a) XRD analysis of the SnSe nanoribbons (76 h, 45 ml grown sample). The two strong peaks can be indexed to phase pure orthorhombic Pnma 62 SnSe. (b) Raman analysis of the same sample as in (a). Peaks are clearly seen, corresponding to the \(A_{1g}, B_{3g}, A_2^g\) and \(A_3^g\) Raman modes of SnSe respectively (peaks fitted with a Lorentzian function). (c) Laser power dependent Raman spectra for SnSe nanoribbons. (d) Position of different Raman modes of SnSe as a function of laser power. Solid lines are linear fit to the experimental data.
potential of SnSe nanoribbons as an efficient thermoelectric material. However, detailed studies of the thermal, electrical and thermoelectric properties of the SnSe nanoribbons are necessary to verify its true potential in thermoelectric devices, which we delegate to a later study.

Figure 4(a) shows an EDX map of a nanoribbon (after 72 h in the 125 ml vessel), revealing the uniform distribution of Sn and Se throughout the nanoribbons, without any elemental segregation. The nanoribbon morphology and thickness were investigated with TEM analysis (figure 4(b)).
however the PVP coating of the nanoribbons presented difficulty with the imaging. A SnSe nanoribbon oriented close to the [001] zone axis is presented in figure 4(b), with a fast Fourier transform (FFT) pattern in the top left inset, indicating the (200) diffraction spot, which is oriented perpendicular to the nanowire growth direction. A thickness of \(~11\) nm was measured, and as this is along the (100) direction, this represents the thickness of the nanoribbon. The bottom right inset of figure 4(b) schematically represents how the nanoribbon is oriented in the figure, highlighting how the measured \(~11\) nm value corresponds to the nanoribbon thickness. A thickness of \(~11\) nm for the nanoribbon would indicate about 19–20 SnSe layers in the nanoribbon structure. The exact layer number is made difficult to determine by the obscuring effect of the PVP present. In general, thicknesses of between \(~11\) and 13 nm were observed from the TEM analysis of the SnSe nanoribbons.

In order to facilitate clearer imaging of the crystal structure of the SnSe nanoribbons they were washed in ethanol to remove the majority of the PVP. Figures 4(c) and (d) show TEM images of ethanol washed SnSe nanoribbons grown after 76 h in the 45 ml vessel, with indexed SAED patterns as insets, orientated along the [100] and [001] zone axes, respectively. Another interesting feature of tailored nanoribbon morphologies is the possibility of defined edge states. The nature of the edge states depends not only on the crystal structure of the system, but also on the way they terminate, which influences the physical properties of the nanoribbons. After removing the PVP, the zigzag structure of SnSe along the (010) direction is clearly seen in the HRTEM image in the inset of figure 4(c). This is of particular interest due to the theoretically predicted semiconductor to metallic transition in very thin Zigzag SnSe nanoribbons [26, 27]. The larger number of morphological variables, such as size, edge-type and thickness, associated with nanoribbon structure over 2D films could allow further scope for better manipulation of their physical properties. After removing the PVP, a dark contrast amorphous layer is also apparent on the surface of the nanoribbons. This amorphous coating is darker in contrast to the previous PVP coating (figure 4(b)) and is shown to consist of SnO\(_2\) by EELS analysis (see supplementary information figure S8). SnSe has been shown to oxidise rapidly in air, forming a SnO\(_2\) surface of \(~6\) nm [12, 53], which is consistent with the thickness of the amorphous layer seen in the TEM images; indicating that this layer is formed post-growth, and post removal of the PVP due to atmospheric surface oxidation. The total thickness of the SnSe nanoribbon, visible when it is viewed down the [001] zone axis in figure 4(d), was seen to be \(~12\) nm, which is also in agreement with the nanoribbon thickness seen when the PVP was present. However, the crystalline core region is reduced to a thickness of \(~6\) nm, with the rest being formed of amorphous SnO\(_2\). This indicates that the PVP appears to offer some protection to atmospheric oxidation, and removal of the PVP should only be performed directly before the implementation of the SnSe nanoribbons into a device. The SnSe nanoribbons are single crystalline throughout their core, as shown in the HRTEM images and corresponding SAED patterns (figures 4(c) and (d) with SAED patterns in insets). The single crystalline nature of these SnSe nanoribbons is important to note, as high yield solution phase growth methods can commonly result in the formation of nanomaterials with a polycrystalline or highly defected crystal structure [34, 54]. Growth directions along the (010) crystal direction of SnSe was observed for every nanoribbon studied (figures 4(c) and (d)), demonstrating a high level of uniformity in the single crystalline nanoribbons. The crystal structure of a (010) oriented SnSe nanoribbon is illustrated in figures 4(e) and (f). Most previous studies on 1D SnSe nanowires have shown growth directions along either the (011) or (100) directions, [12, 39, 41] but in our case, for all nanoribbons analysed, the growth direction was found to be (010). This growth direction for 1D SnSe has only ever been shown for oleylamine [29] and ethanolamine [30] assisted methods, producing short nanorods and large nanobelts respectively. Growth along the (010) direction leads to the exposure of energetically unfavourable high energy surface planes for SnSe [55], demonstrating the necessity of a suitable capping agent. In our approach PVP selectively binds to the (100) and (001) SnSe crystal surfaces, facilitating growth along the (010) direction. Of note, the thermoelectric measurement along the (010) crystal direction has previously been reported to exhibit an ultralow lattice thermal conductivity and a very high thermoelectric figure of merit for bulk SnSe crystals [22]. Utilising (010) oriented SnSe nanoribbons for future thermoelectric studies would thus be of significant interest. The potential of the SnSe nanoribbons in thermoelectric is also demonstrated through temperature dependent Raman spectroscopy (figures 3(c) and (d)).

The growth of SnSe in the 1D nanoribbon structure can be explained through the choice of precursors and PVP as surface passivating agent (figure 5). The growth mechanism of such SnSe flakes and nanoribbons is illustrated in figure 5. The growth of SnSe flakes has previously been shown to initiate as a high surface energy (010) and (001) faceted SnSe flake, with the lower surface energy (011) and (0–11) facets forming the surface planes in the later stages of growth (top row–figure 5) [55]. The basal plane of the layered crystal (100) surface possesses the smallest surface energy, resulting in majority of (100) surface in the nanoribbons. For the formation of SnSe nanoribbons, PVP acts to confine the crystal growth in the early (010) and (001) faceted stage and then restricts the growth in other directions by allowing the adatoms addition preferentially along the (010) plane (illustrated in the bottom row of figure 5).

Capping ligands like PVP plays a significant role in inducing directional growth by selectively adsorbing to specific crystal facets and inhibiting the growth of these facets. PVP reduces the surface free energy of a specific type of facet through selective binding or chemisorption. Without any ligand (PVP in our case), the growth of a nanocrystal will proceed along the path that minimizes the total surface free energy of the system. For SnSe, the absence of any capping molecules leads to the formation of nanosheet morphology, as described in the top row of figure 5. However, ascribed to their unique molecular structures and conformations, surface ligands such as PVP can preferentially bind to a certain crystal facet. This mainly depends on the binding energy which is influenced by the symmetry between the molecule
and crystal surface, the surface-sensitive balance between direct bonding and other kinds of attraction (e.g. van der Waals attraction), etc. The role of PVP in surface selective binding and directional growth of metals (e.g. Ag) is well documented. However, a detailed study, incorporating the density functional theory calculations is required for an accurate understanding of the role of PVP in the directional growth of monochalcogenide based nanostructures such as SnSe nanoribbons and nanowires, which is delegated to a future study.

Due to the layered crystal structure of SnSe, crystal growth tends to dominate along the b and c crystal axes, with minimal growth along the a-axis, upon which the SnSe layers are stacked. Thus the use of SeO2 with a chain structure, and PVP as a passivating agent was crucial for the preferential 1D growth of the layered SnSe structure. The chain structure exhibited by SeO2 is believed to lead to the formation of a chain (Se)n(PVP)m complex, similar to chain (Se)n(Oleylamine)m complexes observed in previous 1D SnSe nanowire growth [29]. To these (Se2−)-PVP seeds, SnCl2.2H2O is then added as a source of Sn2+ which then diffuses into the Se-PVP to form SnSe-PVP nanoparticles (figures S4 and S5 in the supporting information). The SnSe nanoribbons then grow from the SnSe nucleation points, with the aid of PVP as a 1D directing agent. 1D SnSe growth with oleylamine as growth directing reagent at very similar growth temperature (∼175 °C) resulted in the formation of nanorod structure [29]. However, much stronger confinement with long-chain PVP polymer resulted in the formation of a much thinner 1D structure in the form of nanoribbons. PVP is a strong capping agent that interacts with the SeO2 through the strong ionic bonds between the Se2− ions and the amide group oxygen in the PVP chain. Therefore, it is believed that PVP stabilizes the dissolved precursor by amide group steric and electrostatic stabilization and facilitates the growth of nanoribbons. The nanoribbon width and thickness are not seen to increase by any significant amount with increasing length or reaction time, and after the initial nanoribbon formation, the growth continues only by addition along the (010) crystal facets. Additional reaction times lead to further addition along with the (010) crystal facets, and thus longer nanoribbons. The growth of SnSe has commonly been reported to result in the formation of a nanoflake morphology, without the use of any additional surfactants [56].

In a typical growth, SeO2 was heated in ethylene glycol to reduce the Se4+ to Se2− [57], which is important as tin selenides have commonly been reported to form both SnSe and SnSe2 phases, where Sn exists in both 2+ and 4+ oxidation states respectively. The oxidation of Sn2+ to Sn4+ has previously been shown to be suppressed, during the reaction, by the addition of a surfactant such as L-ascorbic acid, aiding in the formation of phase pure SnSe [58]. The XPS spectrum (Se spectrum) of the ethanol washed SnSe nanoribbon sample
Verifies the single oxidation state of Se\(^{2-}\), supporting the high phase purity of the SnSe sample. Though the Sn XPS spectrum (figure 6(b)) reveals three oxidation states for Sn; Sn\(^0\), Sn\(^{+2}\) and Sn\(^{+4}\). This is in agreement with the EELS analysis (figure 6(c)), which reveals that the Sn\(^{+4}\) results from the surface oxidation of the SnSe nanoribbons, with the body of the nanoribbon remaining single crystalline SnSe. The presence of oxygen at the surface region gives rise to the splitting of EEL spectra at 530–540 eV (figure 6(c)), in contrast to no splitting feature shown in the bulk region of nanoribbon, revealing that the amorphous surface layer (region 1) is composed of SnO\(_x\). SnSe has previously been reported to oxidise rapidly in the air [12, 53]. PVP is expected to offer improved protection from this post-growth oxidation, but oxygen diffusion through the PVP layer will still cause small diameter nanostructures to oxidise gradually over time [59].

**Conclusions**

The synthesis of few-layer thick single crystalline SnSe nanoribbons has been successfully achieved through a simple solution-based method, using readily available precursors. Non-toxic PVP was used for the first time to direct the 1D growth of SnSe. The easy to scale-up synthesis approach using non-toxic PVP to grow SnSe nanoribbons is ideal for nanoribbons application in biomedical devices, batteries, solar cells, memory devices and thermoelectric generators. SnSe nanoribbons demonstrated high morphological and structural quality, and promising thermophysical properties. The synthesised nanoribbons all grow along the \(\langle 010 \rangle\) direction, which has been previously reported to possess a remarkably low lattice thermal conductivity along with this crystal orientation, highlighting the significance of these nanoribbons for potential applications in future thermoelectric devices. The growth along the \(\langle 010 \rangle\) direction is uncommon as the crystal
growth along this direction exposes the high surface energy planes in the structure. The uncommon growth along the (010) direction is achieved for SnSe nanoribbons through the selective binding of PVP to the (100) and (001) surface planes. A better understanding of the nature of the influence of PVP in SnSe systems could lead to future possibilities in material design and non-toxic growth strategies for the solution phase synthesis of SnSe and other group IV monochalcogenides. The growth of very thin (010) oriented SnSe nanoribbons has not previously been reported, and the successful synthesis of SnSe nanoribbons also enables the future exploration of the previously theorised novel properties of SnSe nanoribbons and other nanoribbon structures, such as a width-dependant tuneable bandgap for application in electronic, energy, and memory devices.

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Data availability statement
The data that support the findings of this study are available upon reasonable request from the authors.

Conflict of interest
The authors declare no competing financial interest.

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