

Controlled Crystal Growth of Indium Selenide, In_2Se_3 , and the Crystal Structures of $\alpha\text{-In}_2\text{Se}_3$

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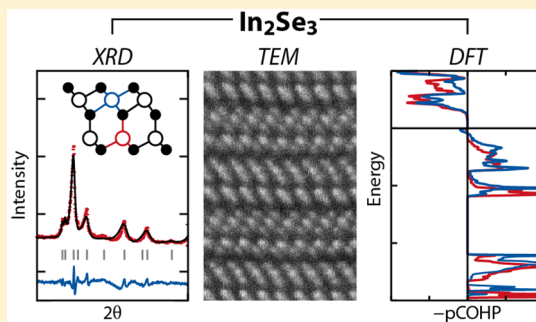
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S Supporting Information

ABSTRACT: In_2Se_3 has been known for over 100 years and recently attracted interest as a promising candidate for a variety of applications, such as solar cells, photodiodes, and phase-change memories. Despite the broad concern for possible uses, its polymorphism and structure are poorly characterized. By combining X-ray diffraction, transmission electron microscopy, and quantum-chemical calculations, we present here the crystal structures of two layered room-temperature polytypes: 3R and 2H In_2Se_3 . Both polymorphs are stacking variants of the same Se–In–Se–In–Se layers comprising two coordination environments for the In atoms, one tetrahedral and one octahedral. By using chemical-bonding analysis, we look at the different In positions in $\alpha\text{-In}_2\text{Se}_3$ and compare them to those in the metastable β -phase.



INTRODUCTION

The archetypal phase In_2Se_3 is known for its outstanding photoelectric properties and possible applications in solar cells^{1,2} or photodiodes.³ Moreover, In_2Se_3 shows a plethora of other promising properties like thermo^{4–6} and ferroelectricity,^{7–9} as well as a strong contrast in electrical properties between its different phases. Recently, the phase therefore gained interest as a phase-change material (PCM) for data storage.^{10–12} Conventional PCMs switch between an amorphous and a crystalline state,¹³ requiring the melting and also quenching of the material, which results in high stresses within the material and poor energy efficiency for the device. To achieve better performance, recent studies suggest the use of two different crystalline phases to store information. For layered compounds these unconventional PCMs have been dubbed interfacial PCMs (iPCM).¹⁴

Although In_2Se_3 is currently being discussed as a promising candidate for many applications, its crystal structures and the crystallization behavior of its polymorphs remain unclear, at best. Therefore, a thorough structural characterization of In_2Se_3 is of high interest, because it enables us to understand its physical properties and allows for the intelligent design of new data storage devices.

A compound with the In_2Se_3 stoichiometry has already been known since 1910¹⁵ and was characterized by Klemm in

1934.¹⁶ The polymorphism and phase diagram of In_2Se_3 are controversially discussed in the literature.^{17–21} There are supposed to be at least four different polymorphs in the bulk material at ambient pressure: α , β , γ , and δ . In addition, $\alpha\text{-In}_2\text{Se}_3$ seemingly consists of two different stacking variants, the so-called 2H and 3R phases. High-pressure phase transitions starting from 3R In_2Se_3 were investigated very recently.²² Concerning the crystal structures of $\alpha\text{-In}_2\text{Se}_3$, lots of different structure models are found in the literature,^{4,9,23} where recently a model for the 2H phase was derived from electron microscopy,²⁴ but atomic parameters are, to the best of our knowledge, unknown.

In this work, we present, for the first time ever, a detailed structural characterization of the two $\alpha\text{-In}_2\text{Se}_3$ variants. The structural characterization using single-crystal and powder X-ray diffraction is complemented by high-resolution scanning transmission electron microscopy (HR STEM), as well as density-functional theory (DFT) calculations.

METHODS

Synthesis. Samples of 3R In_2Se_3 were synthesized from the elements (In, HMW Hauner 99.999%; Se, MaTeck 99.999%) in

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evacuated quartz ampules. Samples were first heated to 500 °C within 1 h and then heated to 900 °C over the course of 20 h to prevent Se from evaporating too fast. The product was then held at 900 °C for 20 h and quickly cooled to room temperature.

Chemical transport^{25,26} of In₂Se₃, using a temperature gradient from 800 to 700 °C and iodine (Merck 99.5%) as transport agent, resulted in very thin and brittle plate-like crystals of 2H In₂Se₃. Unfortunately, we were unable to isolate a single-crystalline specimen of the 2H crystals since they are mechanically very weak, and trying to cut or break larger crystals resulted in polycrystalline pieces.

Characterization. A sample tempered at 250 °C for 160 h and quenched to room temperature in air provided a twinned specimen of 3R In₂Se₃ with approximate dimensions of 0.05 × 0.05 × 0.01 mm³. It was selected for a single-crystal diffraction experiment at room temperature, where 2018 reflections were collected in ω mode on a Bruker D8 goniometer equipped with an APEX CCD detector and an Incoatec microsource (Mo K α radiation, λ = 0.71073 Å). Crystallographic and structure refinement data are shown in Table 1. The

Table 1. Crystallographic and Structure Refinement Data of 3R In₂Se₃

formula	In ₂ Se ₃
formula weight (g/mol)	466.52
space group	R3m (No. 160)
<i>a</i> (Å)	4.026(4)
<i>c</i> (Å)	28.750(11)
<i>V</i> (Å ³)	403.6(8)
<i>Z</i>	3
diffractometer	Bruker D8 goniometer with APEX CCD detector
Θ range	2.125–28.602
<i>hkl</i> ranges	−4 ≤ <i>h</i> ≤ 4; −5 ≤ <i>k</i> ≤ 5; −37 ≤ <i>l</i> ≤ 37
no. reflns	2018
no. indep reflns	310
<i>R</i> _{int}	0.0639
no. params	20
<i>R</i> ₁ ; <i>wR</i> ₂ (all data) ^a	0.1084; 0.2444
GOft ^a	1.097
diffraction peak and hole (e Å ^{−3}) ^a	5.472/−8.642
CCDC number	1855522

^aHigh residual parameters are explained in the text.

diffraction pattern was interpreted with the help of CELL_NOW:²⁷ 346 out of a subset of 379 reflections could be indexed when two domains with the lattice parameters for 3R In₂Se₃ known from powder diffraction were taken into account. The relationship between these domains matched the expectation for obverse/reverse twinning with the matrix ($\bar{1}00$; $0\bar{1}0$; 001). After integration with SAINT+, an absorption correction with TWINABS²⁸ based on multiple scans was applied. The structure solution by direct methods²⁹ was in full accord with the atomic coordinates as derived from powder diffraction. When only the major domain was considered, refinement with SHELXL³⁰ converged to residual values of *wR*₂ = 0.44 (all data) and *R*₁ = 0.20 and left a residual electron density of 32 e Å^{−3}. When obverse/reverse and inversion twinning in the noncentrosymmetric space group R3m (No. 160) was taken into account, these agreement factors improved to *wR*₂ = 0.24 (all data) and *R*₁ = 0.10, and the residual electron density decreased to 5 e Å^{−3}. In refining these intensity data, the more precise lattice parameters from the powder experiments were used.

Although the aforementioned domains could be assigned unambiguously, they cannot completely describe the diffraction pattern. This limitation is reflected in unsatisfactory residual values and the rather high residual electron-density maxima and minima. Together with In₂, the most pronounced electron-density fluctuations in a final difference Fourier map are situated along the crystallographic *c* axis, at fractional coordinates of 0, 0, 0.140 (5.4 e Å^{−3}) and 0, 0,

0.074 (−8.6 e Å^{−3}). A view of these most relevant electron-density maxima and minima is provided in the Supporting Information. Less pronounced local electron-density fluctuations are encountered in the vicinity (0.6–0.9 Å) of the heavy atoms.

Tentative inclusion of additional smaller domains of the same compound caused less stable data integration and did not improve the refinement results. Additional details concerning the structure determination for 3R In₂Se₃ are available in CIF format and have been deposited under the CCDC entry number 1855522. Copies of the data can be obtained free of charge from CCDC.

The polycrystalline samples of 3R In₂Se₃ and crystals of 2H In₂Se₃ were mortared and measured by powder X-ray diffraction using a STADI MP diffractometer (STOE, Darmstadt, Germany) with monochromatic Mo K α radiation and a PSD detector. The intensity data were refined using the Fullprof program package.^{31,32} Crystallographic and Rietveld refinement data are shown in Table 2. To

Table 2. Rietveld Refinements of Powder XRD Data

	2H α -In ₂ Se ₃	3R α -In ₂ Se ₃
space group	<i>P</i> 6 ₃ <i>mc</i> (No. 186)	<i>R</i> 3m (No. 160)
<i>a</i> (Å)	4.023(4)	4.026(4)
<i>c</i> (Å)	19.217(11)	28.750(11)
<i>V</i> (Å ³)	269.4(2)	403.6(8)
<i>Z</i>	2	3
χ^2	2.86	1.76
<i>R</i> _{Bragg}	8.25	11.0
<i>R</i> _F	8.47	8.12
<i>R</i> _p (%) ^a	1.53	2.30
<i>wR</i> _p (%) ^a	2.31	3.27

^aNot corrected for background.

account for anisotropic broadening due to two-dimensional defects that could also be seen in the STEM images (see Figure S1), an anisotropic size model was employed while refining the diffraction pattern. For 3R In₂Se₃, the single-crystal data were used as a starting model with fixed atom sites and displacement parameters. For 2H In₂Se₃, due to a lack of single-crystal data, a starting model with quantum-chemically optimized atomic positions was used to refine the diffraction pattern. High-resolution XRD measurements at a synchrotron beamline are planned to fully refine the crystal structure. Both powder patterns show a reflection at 13.7° in 2 Θ that might be indexed by β -In₂Se₃.

For TEM measurements, lamellas were produced by focused ion beam (FIB) technique employing an FEI Dual Beam Helios NanoLab system. The lamellas were cut along the *c* axis of In₂Se₃ single crystals.

The high-angle annular dark-field scanning transmission electron microscopy (HAADF-STEM) imaging together with STEM-energy dispersive X-ray spectroscopy (STEM-EDX) mappings were performed on an FEI Titan electron microscope, equipped with a Cs-aberration corrector for the probe-forming lens operated at 200/300 kV acceleration voltage. For EDX mapping a “Super-X” wide solid angle EDX detector was used at a voltage of 200 kV.

Computational Details. Density-functional theory (DFT) calculations employed the “D3” van der Waals correction³³ with dampening³⁴ on top of the PBE functional.³⁵ This approach reproduces structural parameters well and accounts for dispersion interactions between quintuple-layer blocks. We used plane-wave basis sets and the projector augmented wave (PAW) method³⁶ as implemented in the Vienna *Ab Initio* Simulation Package (VASP).^{37–39} The energy cutoff for the plane-wave expansion was set to 500 eV with an electronic convergence criterion of 10^{−7} eV. Structural optimization was performed until residual forces fell below 1 × 10^{−3} eV Å^{−1}. Reciprocal space was sampled on Γ -centered *k*-point grids with densities between 0.01 and 0.03 Å^{−1} for structural relaxation. Chemical-bonding analyses of plane-wave data by Crystal Orbital Hamilton Population (COHP)^{40,41} as well as Mulliken charge

analysis were done using LOBSTER.^{42–44} The following pseudopotentials have been employed: In, 4d, 5s, 5p; and Se, 4s, 4p.

RESULTS

Figure 1 shows an HR HAADF-STEM image of two different polycrystalline In_2Se_3 samples. Because the intensity in

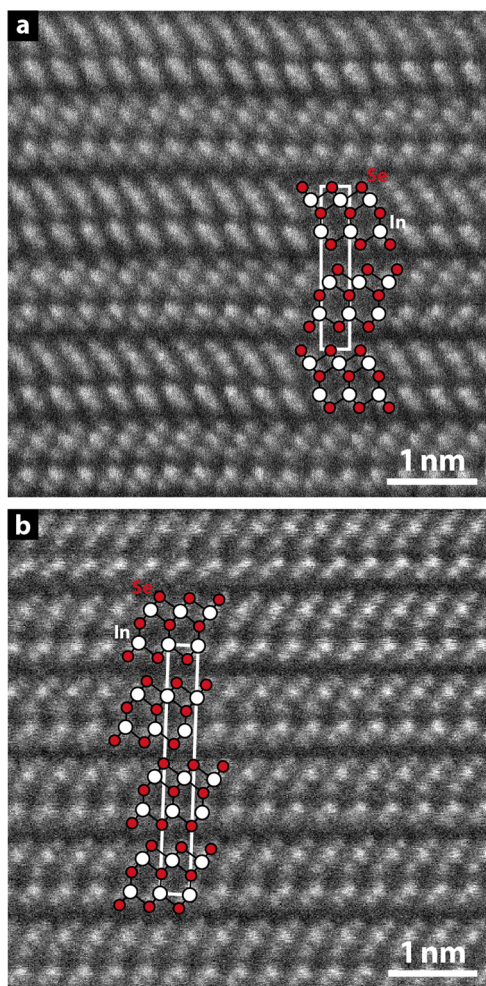


Figure 1. High-resolution HAADF-STEM Z-contrast images of (a) 2H and (b) 3R In_2Se_3 together with the structural models.

HAADF-STEM is roughly proportional to $Z^{1.8}$, this allows the straightforward interpretation of the images: In-rich columns appear slightly brighter than Se columns ($Z_{\text{In}} = 49$, $Z_{\text{Se}} = 34$). The results are confirmed by STEM-EDX mapping as shown in Figure S2. Thus, both samples are arranged in a layered structure of quintuple Se–In–Se–In–Se layer blocks stacked in one direction. The blocks are separated by van der Waals gaps.

The two In atoms in the layers are coordinated differently, one tetrahedrally and the other octahedrally. The difference between 2H and 3R In_2Se_3 lies in the stacking of these layers. The 2H and 3R In_2Se_3 structure models as created to fit the HAADF-STEM data were further optimized by quantum-chemical calculations and then used for refining powder XRD data. The 3R In_2Se_3 structure was confirmed by single-crystal diffraction.

Due to its layered van der Waals-like crystal structure, α - In_2Se_3 shows a large number of two-dimensional defects perpendicular to its stacking in the bulk phase. This results in

strong anisotropic broadening and preferred orientations in the PXRD data as visible in Figure 2.

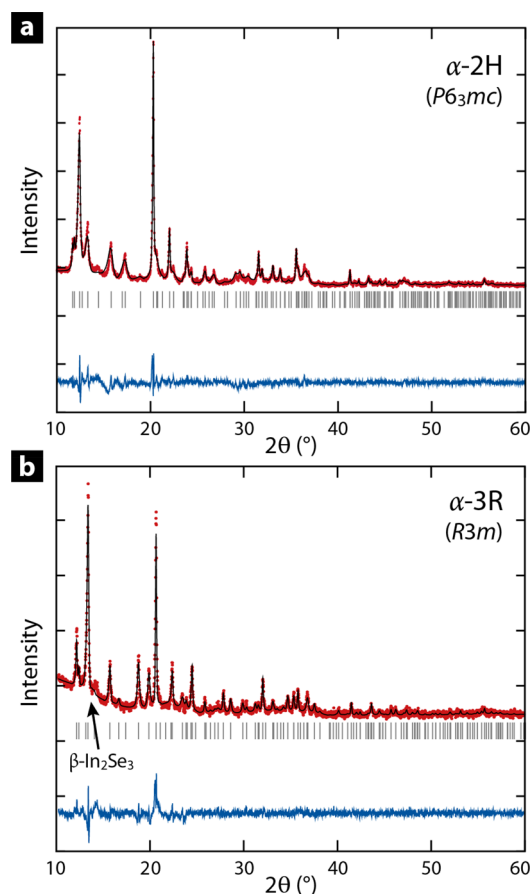


Figure 2. Refined X-ray powder diffraction patterns of (a) α -2H and (b) α -3R In_2Se_3 samples.

2H In_2Se_3 shows hexagonal symmetry with space group $P6_3mc$ and five different atomic sites (In1 on 2a, In2 on 2b, Se1 on 2a, Se2 on 2b, Se3 on 2a). Popovic et al. postulated space group $P6_3/mmc$ but without a structure model suggestion.¹⁹ Each layer consists of two indium atoms with different coordination polyhedra: In1 is coordinated tetrahedrally by four Se atoms while In2 is octahedrally coordinated by six Se. Moving further, the 3R structure consists of identical quintuple layers in a different stacking. Instead of the “zigzag” motif of the 2H structure (see Figure 1a), the layers differ only on the basis of a translation along the ab plane. In 3R, five atomic sites (In1, In2, Se1, Se2, Se3; all on 3a positions) exist, arranged in a rhombohedral ($R3m$) crystal structure. By turning and moving the layers along the ab plane, both structures are easily transferred into each other.

A similar structure model for the 3R structure exists since 1966 and goes back to Osamura et al.,⁴⁵ who already described a quintuple-layer structure with InSe_4 tetrahedra and InSe_6 octahedra. The interatomic distances in their structure model, however, seem questionable, for example, homoatomic Se–Se distances of 2.61 Å in comparison to a heteroatomic In–Se distance of 3.51 Å. These questionable details by Osamura et al. probably go back to a simple printing mistake because their structure is close to ours if one exchanges the z parameters of Se2 and In1. Ye et al. suggested another layered structure with all tetrahedrally coordinated indium atoms.⁴⁶ However,

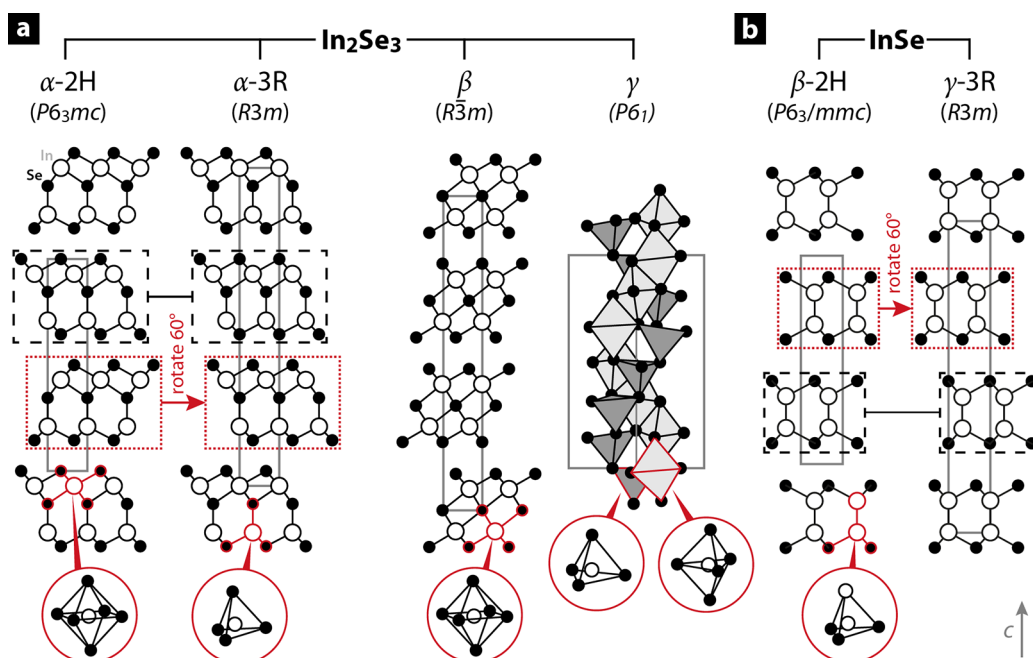


Figure 3. Projection of (a) α -, β -, and γ - In_2Se_3 crystal structures and (b) stacking variants of α -InSe along c , highlighting different coordination environments.

arranging quintuple Se–In–Se–In–Se layers with In atoms coordinated only tetrahedrally will result in some single-coordinated Se atoms. This is highly questionable already from chemical intuition, and DFT calculations show an instability of the all-tetrahedral coordination toward one octahedrally coordinated In atom. Almost 50 years later, in 2015, Debbichi et al.²³ postulated a new structure model for 3R In_2Se_3 by quantum-chemical calculations, which is isostructural to the structure found by our group. Very recently, Zhou et al. could experimentally confirm this very structure by electron microscopy of In_2Se_3 nanosheets.⁹

For 2H In_2Se_3 , one structure model was already postulated by Semiletov in 1961,⁴⁷ also with all tetrahedrally coordinated In atoms. Like the 3R model from Ye et al., this model suffers from the single-coordinated selenium atoms. As mentioned before, however, a better model has recently been derived from electron microscopy results by Xue et al.²⁴

Above 200 °C, In_2Se_3 is supposed to undergo a phase transition into β - In_2Se_3 , which crystallizes in the Bi_2Te_3 structure type. This phase also consists of Se–In–Se–In–Se quintuple layers, but here, In is octahedrally coordinated throughout. β - In_2Se_3 is stable in nanosheets at room temperature⁴⁸ and was reported to be rapidly quenchable to room temperature as a bulk phase.¹⁹ However, we were unable to reproduce this behavior.

Figure 3 displays the structural relationships between α -, β -, and γ - In_2Se_3 as well as two stacking variants of indium monoselenide (InSe). The crystal structure of γ - In_2Se_3 is well-known and was solved by single-crystal refinement.⁴⁹ It crystallizes in a defect wurtzite type with vacancy spirals. This is the only known no-layered polymorph of In_2Se_3 and consist of tetrahedrally and pentagonally coordinated In atoms. In addition, there are two more potentially layered structures of In_2Se_3 reported in the literature: κ - In_2Se_3 crystallizes in nanosheets only and is supposed to be similar to the α -phases,^{50,51} while δ - In_2Se_3 is a high-temperature phase stable above 720 °C¹⁹ whose crystal structure is still unknown.

Indium monoselenide (InSe) also crystallizes in a layered structure but with quadruple Se–In–In–Se layers instead. Surprisingly, the polymorphism of InSe shows exactly the same stacking behavior as α - In_2Se_3 : one hexagonal stacking (β -InSe, $P6_3/mmc$), where every second layer is rotated by 60° around [001], and one rhombohedral stacking (γ -InSe, $R3m$), arranged by translation within the ab plane.

Heterointeratomic distances in the two α -phases are very similar and range from 2.55 to 2.64 Å for the tetrahedrally coordinated In atom. For the octahedrally coordinated indium, the Se–In distances are larger, as expected (2.71–2.85 Å). These distances are in excellent agreement with those found in other compounds of the In–Se phase diagram, with similar coordination polyhedra. In γ - In_2Se_3 one In atom is also tetrahedrally coordinated with interatomic distances between 2.57 and 2.63 Å.⁴⁹ An InSe_6 -octahedron can be found in the In_6Se_7 crystal structure, with In–Se distances of 2.65–2.89 Å.⁵²

Given this overview of its structures, one may well ask why In_2Se_3 prefers a mixed coordination instead of an all-octahedral one. To answer this question, we performed bonding and charge analysis as will be discussed in the following.

Figure 4 shows Mulliken charges and projected COHP plots for α -3R In_2Se_3 and β - and γ - In_2Se_3 . The cationic indium charge within the structures varies from +0.40 to +0.52 e, with the β -phase being the least ionic of the three, while the α -phase shows a slightly stronger cationic behavior of indium in the tetrahedral coordination. Interestingly, most electrons are transferred in the case of γ - In_2Se_3 where indium (within the strongly distorted tetrahedral or trigonal bipyramidal environments) has a charge of +0.50 to +0.52 e. Regarding the question from above, the ionic contributions of tetrahedrally coordinated In atoms are stabilizing the α -phase when compared to the purely octahedral β - In_2Se_3 .

Transitioning from ionic to covalent contributions we show COHP plots in Figure 4b. Here, stabilizing covalent interactions can be seen for the tetrahedrally coordinated In

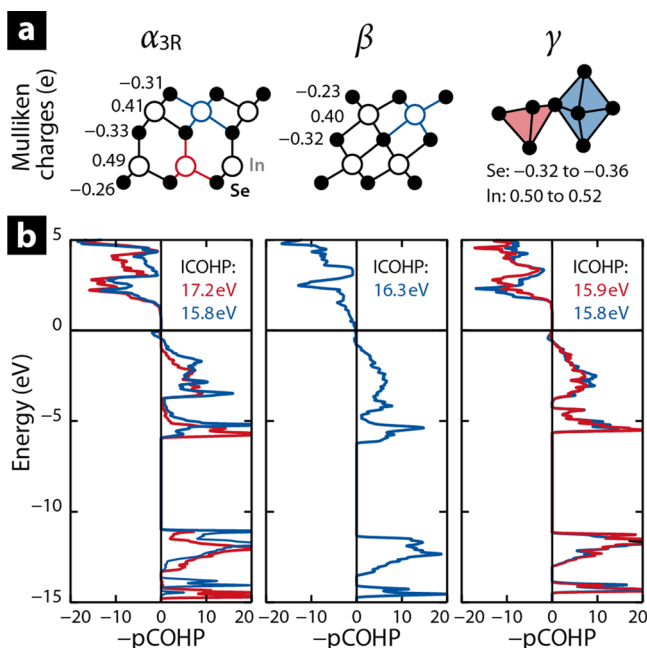


Figure 4. (a) Mulliken charges and (b) chemical-bonding analysis of selected In_2Se_3 polymorphs with bonding levels to the right and antibonding ones to the left. The colors of the COHP plot correspond to the bonds highlighted in the structural fragments shown above. Projected density of states (DOS) plots are provided in the Supporting Information.

atoms (in red) in the α - and γ -phase. For such species, we observe no significant antibonding contributions below the Fermi level, which is contrasted by small antibonding contributions for the octahedrally coordinated atoms in the α - and β -phase. Such antibonding regions are sometimes observed in chalcogenides used as PCMs⁵³ but are overall destabilizing and should be minimized. Furthermore, the integrated COHP values of 3R In_2Se_3 show us that the tetrahedral coordination is indeed favored by covalent interactions as analyzed within the COHP framework.

Hence, indium atoms within In_2Se_3 prefer a tetrahedral environment to an octahedral one both from a covalent as well as an ionic perspective. A purely tetrahedral environment, however, is not possible due to the stoichiometry of the compound, such that there results a proper mix of tetrahedral and octahedral sites as observed in diffraction and electron microscopy.

CONCLUSION

By combining HR-HAADF STEM Z-contrast imaging, single-crystal and powder XRD, and also quantum-chemical calculations, we were able to identify the crystal structures of 2H and 3R In_2Se_3 . These α -phases of In_2Se_3 consist of quintuple Se–In–Se–In–Se layers with two distinct coordination environments for In, one tetrahedral and one octahedral. The two phases are stacking variants, which are selectively accessible by solid-state synthesis. The stability of tetrahedrally coordinated In atoms was elucidated using chemical-bonding analysis tools, which showed an increase in stability for covalent as well as ionic contributions when compared to β - In_2Se_3 with purely octahedral coordination.

ASSOCIATED CONTENT

Supporting Information

The Supporting Information is available free of charge on the ACS Publications website at DOI: 10.1021/acs.inorgchem.8b01950.

Atomic position for 2H and 3R In_2Se_3 , HAADF-STEM Z-contrast image of 3R In_2Se_3 , EDX mapping of Se and In in 2H In_2Se_3 , location of the residual electron densities from single-crystal refinement, optical micrographs of polycrystalline samples of 3R and 2H In_2Se_3 , and projected DOS of α_{3R} , β , and γ - In_2Se_3 (PDF)

Accession Codes

CCDC 1855522 contains the supplementary crystallographic data for this paper. These data can be obtained free of charge via www.ccdc.cam.ac.uk/data_request/cif, or by emailing data_request@ccdc.cam.ac.uk, or by contacting The Cambridge Crystallographic Data Centre, 12 Union Road, Cambridge CB2 1EZ, UK; fax: +44 1223 336033.

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Notes

The authors declare no competing financial interest.

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