Tunable spin-orbit coupling in two-dimensional InSe

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We demonstrate that spin-orbit coupling (SOC) strength for electrons near the conduction band edge in few-layer γ-InSe films can be tuned over a wide range. This tunability is the result of a competition between film-thickness-dependent intrinsic and electric-field-induced SOC, potentially, allowing for electrically switchable spintronic devices. Using a hybrid k·p tight-binding model, fully parameterized with the help of density functional theory computations, we quantify SOC strength for various geometries of InSe-based field-effect transistors. The theoretically computed SOC strengths are compared with the results of weak antilocalization measurements on dual-gated multilayer InSe films, interpreted in terms of Dyakonov-Perel spin relaxation due to SOC, showing a good agreement between theory and experiment.

I. INTRODUCTION

Indium selenide (InSe) is a layered semiconductor with already demonstrated high mobility and versatile optical properties1–8. Atomically thin InSe films (exfoliated from bulk crystals9 or produced by chemical vapour deposition10) have already been used to fabricate field-effect transistors (FET devices). Moreover, the persistence of high mobility11–13 of electrons in n-type doped γ-InSe to only few atomic layers3,4,9 in thickness makes it feasible to implement InSe in spintronic devices14. In contrast to the conventional InAs15 or HgTe16 quantum wells, in γ-InSe it is possible to completely suppress the conduction band SOC using applied displacement field, allowing for exceptionally tunable spintronic devices. One of the methods to control the electron spin in semiconductors is to manipulate its spin-orbit coupling (SOC)17–23, and, in this paper, we study the dependence of SOC for two dimensional (2D) electrons near the conduction band edge of InSe films on the number of layers and on the gate-controlled electrostatic doping in the films implemented in the FET geometry24–31.

Below, we use the earlier developed hybrid k·p tight-binding (HkpTB) model for InSe33,34, taking into account the s and pz orbital composition of the lowest conduction subband and self-consistent analysis of the electrostatic potential on each layer35, and show that the dominant term in the SOC in γ-stacked InSe multilayer thin film (any number of layers) has the generic form,

\[ \hat{H}_{SOC} = \alpha (s \times k) \cdot \hat{z}. \] (1)

This is the only linear in wavevector k=(kx, ky) of electron (in the vicinity of the Γ-point) term allowed by C3v point-group symmetry of the lattice of γ-stacked multilayer (the next term in the k·p theory expansion would be of the third order36 in k, hence, much weaker for a
Here, $\chi$ is a layer-number-dependent factor,

$$\alpha(\mathcal{E}_z, N) \approx \alpha_\infty \left( 1 - \frac{\chi}{(N + 2.84)^2} \right) \pm \mathcal{E}_z N. \quad (2)$$

Here, $\alpha_\infty \approx 34.5$ meVÅ is the value of SOC at the conduction band edge of 3D bulk $\gamma$-InSe, $N$ is the number of layers in a thin film, $\chi \approx 14.9$ accounts for the non-linear dependence of bulk SOC on the out-of-plane momentum $k_z$ counted from the bulk A-point band edge, at $k_A = \frac{\pi}{a}$. Also $\mathcal{E}_z$ is the electric field piercing the film, and parameter $N$ quantifies the dependence on the electric field, as shown in the inset of Fig. 1.

The overall strength of SOC in Eq. (2) is determined by the interplay between the intrinsic lattice asymmetry of the crystal (known as Dresselhaus contribution\textsuperscript{41}) and the electric-field-induced symmetry breaking (the so-called Bychkov-Rashba term\textsuperscript{42}). This interplay allows for the tunability of the SOC value, both by choosing the film thickness ($Nd$), and by applying a displacement field in the double-gated (both top- and bottom-gated) devices. The results of our analysis are exemplified in Fig. 1, indicating that a vertically applied electric field $\mathcal{E}_z \sim 0.15-0.20$ Vnm\textsuperscript{-1} would be enough to switch SOC off and on, opening new avenues towards the design of spintronic devices. This form of SOC in a film is the result of $k \cdot \mathbf{p}$ and tight-binding model analysis\textsuperscript{43}, parameterized using density functional theory (DFT) computations of the band structure. The theoretically calculated SOC size was compared with the values of SOC strength extracted from weak antilocalization magnetoresistance, measured in a FET based on a six-layer InSe device. We find a good agreement between theory and experiment in the available range of device parameters.

Below, the paper is organized as follows. In Section II, we compute the SOC coefficient in the lowest conduction subband of InSe using DFT \textit{ab initio} calculations, in Section III, we perform simple perturbative calculations of SOC strength in the lowest conduction subband of bilayer InSe and, in Section IV, we generalise the bilayer formalism for an arbitrary number of layers. Finally, in Section V, we compare the theoretically obtained SOC coefficient with the values experimentally measured in an available InSe-based FET device.

### II. FIRST PRINCIPLES CALCULATIONS OF InSe PARAMETERS

As a background to the hybrid $k \cdot \mathbf{p}$ tight-binding (HkpTB) model presented in this manuscript, we overview the density functional theory band structure of monolayer and few-layer InSe.

Monolayer InSe has pairs of vertically aligned metal

| $E_{v}$ | $t_{vv}^I$ | $t_{cc}^I$ | $E_{v2}$ | $m_{cc}$ | $m_0$ |
|--------|----------|----------|---------|--------|--------|
| -2.79 eV | -0.41 eV | 0.34 eV | -3.5 eV | 0.266 m0 |

TABLE I. (Top) Two-band hybrid $k \cdot \mathbf{p}$ tight-binding parameters extracted from the 14-band model in the bottom table. (Bottom) Hybrid $k \cdot \mathbf{p}$ tight-binding model parameters used in the perturbation theory analysis. Numerical indices in the $b$ and $\lambda$ terms label the symmetry group shown in the character table in Fig. 4. The magnitude of the out-of-plane dipole moments $d_{ce}$, $d_{ve}$, and $d_{vc}$ were obtained from the tight-binding model developed in Ref. 33. The SOC parameter $\lambda_{ce}$ was calculated from the fits performed in Appendix A and the interlayer distance $a_z = 8.32$ Å was obtained from the experimental measurements shown in Ref. 40.

![Fig. 2. Plane-averaged electrostatic potential accounting for ionic and Hartree contributions in a double-bilayer InSe supercell (supercell structure shown as inset).](image)
FIG. 3. DFT-calculated conduction band spin-orbit splitting for mono-, bi-, and tri-layer InSe in for small \( k_x \) near the \( \Gamma \)-point. The finite thickness of the film discretizes \( k_x \) while \( k_x \) and \( k_y \) remain continuous variables. While the cubic Dresselhaus SOC splitting is expected to be zero in the \( \Gamma - M \) direction, in the \((k_x, 0)\) orientation a finite contribution is expected. In contrast to the orientation-dependent cubic SOC, the expected form of the linear SOC splitting (see Eq. (1)) makes this contribution isotropic in \( k \). The shaded region labels the range in \( k_x \) below the Fermi level of a device doped with a carrier density of \( n_c \approx 2 \times 10^{12} \text{cm}^{-2} \). The clear linear spin splitting with \( k_x \) indicates the dominance of the linear SOC terms near the Brillouin zone (BZ) center. Also plotted using stars connected by dashed lines are values of splitting for a monolayer InSe/monolayer hBN heterostructure for three different in-plane stacking configurations. (Inset) Hexagonal BZ of monolayer InSe. The red circle indicates the region in the BZ with wavevector magnitude in the range presented.

atoms in the middle sublayers and chalcogens in the outer sublayers, arranged on a plane into a honeycomb structure. Such a lattice has a \( D_{3h} \) point-group symmetry which includes mirror plane symmetry, rotations by 120°, but not inversion symmetry. In any few-layer \( \gamma \)-InSe film, the \( z \rightarrow -z \) mirror symmetry is broken. This opens a possibility for a weak “ferroelectric” charge transfer between the layers due to layer-asymmetric hybridization between the conduction and valence bands and the resulting built-in electric field in the film which may be relevant for the self-consistent analysis of the on-layer potential in a film with a finite thickness. To find out whether this is of relevance for InSe, or not, we carry out DFT calculations on a supercell with a large vacuum separating two mirror reflected images of a \( \gamma \)-InSe bilayer, to satisfy periodic boundary conditions without affecting the mismatch between vacuum potentials, produced by the double-charge layer due to the charge transfer (see Fig. 2 and inset). For the DFT calculations, we used the generalized gradient approximation (GGA) of Perdew, Burke and Ernzerhof\textsuperscript{44}, with an 12 \( \times \) 12 \( \times \) 1 \( k \)-point grid and a plane-wave cutoff energy of 600 eV, implemented in the VASP code\textsuperscript{45}. Monolayer atomic structure parameters, and interlayer distances, are taken from an experimental reference for the bulk crystal\textsuperscript{40}. We find that the charge transfer between the layers is small, yielding a \( \approx 2 \) meV vacuum potential difference across the bilayer in Fig. 2, which is so small that it will be neglected for the rest of the manuscript.

Due to its mirror symmetry, the monolayer Hamiltonian cannot include \( s_x \), and \( s_y \) operators, that is, it does not display a 2D SOC. However, its symmetry allows for spin-orbit splitting in the form of\textsuperscript{41,46}

\[
\hat{H}_{so} = \gamma k^3 \sin(3\phi) \hat{s}_z
\]

where \( \phi \) is the polar angle with respect to the \( \Gamma - M \) direction and \( \hat{s}_z \) is the third Pauli matrix. This is reflected by the results of DFT computations of conduction band dispersion in mono-, bi-, and trilayers shown in Fig. 3(a).

Note that the spin polarization of the computed states is in \( z \)-direction only for monolayers, whereas for bi- and trilayers, where it has a linear dependence announced in Eq. (1), it reflects in-plane spin splitting. In fact, for the
range of in-plane wavenumbers corresponding to feasible doping densities, the spin splitting in the monolayer is negligibly small.\cite{47} We also carried out DFT calculations for a heterobilayer consisting of monolayer of InSe, and monolayer of hBN (the latter was strained to give commensurability with a lattice constant \(a_{\text{hBN}} = a_{\text{InSe}}/\sqrt{3}\) and rotated to align the armchair direction of hBN with the zigzag direction of the InSe). We take the interlayer distance as 0.333 nm between the middle of hBN and the nearest plane of Se atoms. A dipole correction was applied, and we considered three in-plane configurations: (1) boron directly above indium, (2) nitrogen above indium (the hBN is inverted in-plane), and (3) configuration (1) with the hBN shifted in-plane by half the B-N vector. The spin-orbit splitting near \(\Gamma\) in the (InSe-dominated) conduction band edge is plotted for all 3 configurations in Fig. 3. For the monolayer InSe/monolayer hBN heterostructure, we obtain a SOC which depends very weakly on the configuration, with a strength similar to that of the isolated InSe bilayer.

### III. SPIN-ORBIT COUPLING IN InSe BILAYER

InSe belongs to the family of group-III metal-monochalcogenides with the \(s\) and \(p\) orbitals of In and Se dominating the low-energy dispersion in the vicinity of the \(\Gamma\)-point.\cite{33,43,46} In the bottommost conduction band \(c\) and in the topmost valence band \(v\), the atomic orbital composition is mainly dominated by the \(s\) and \(p_z\) orbitals of both In and Se. The deeper valence bands \(v_1\) and \(v_2\) are prominently Se \(p_x\) and \(p_y\) orbitals which are naturally split by the atomic SOC of the Se atoms.

In going from monolayer to bilayer \(\gamma\)-stacked InSe (see Fig. 5), the mirror plane symmetry is broken, reducing the symmetry from the point-group \(D_{3h}\) to \(C_{3v}\). This allows for a linear in momentum SOC splitting in the form presented in Eq. (1) prescribed by the third-order rotation symmetry axis.\cite{48} Consequently, the interlayer hoppings need to account for the reduction of the global symmetries of the bilayer, leading to a finite Dresselhaus SOC. This appears via the interlayer mixing of the opposite \(z\)-parity bands.

We construct a bilayer Hamiltonian using monolayer Hamiltonians described in Ref. 33 taking into account interlayer hopping and the intralayer interband spin-orbit coupling:\cite{49}

\[
\hat{H} = \hat{H}^{(0)} + \delta \hat{H} = \begin{pmatrix}
H_{11}^{(0)} + \delta H_{11} & \delta H_{12} \\
\delta H_{21} & H_{22}^{(0)} + \delta H_{22}
\end{pmatrix}, \quad (4)
\]

For the analysis of SOC in the bilayer, band edge states in the constituent monolayers,

\[
\Psi^T \equiv [c_1^\dagger, c_2^\dagger, v_1^\dagger, v_2^\dagger, v_1^\dagger p_x, v_1^\dagger p_y, v_2^\dagger p_x, v_2^\dagger p_y, v_1^{\dagger p_x}, v_1^{\dagger p_y}, v_2^{\dagger p_x}, v_2^{\dagger p_y}],
\]

for the bands described in Section II, will be characterised by their respective band energies, neglecting an almost parabolic band dispersion.

\[
\hat{H}^{(0)}_{11(22)} = \begin{pmatrix}
-U_{1(2)} & 0 & 0 & 0 & 0 & 0 \\
0 & E_{c_1} - U_{1(2)} & 0 & 0 & 0 & 0 \\
0 & 0 & E_v - U_{1(2)} & 0 & 0 & 0 \\
0 & 0 & 0 & (E_{c_1} - U_{1(2)})\hat{I}_v & 0 & (E_{c_2} - U_{1(2)})\hat{I}_v \\
0 & 0 & 0 & 0 & (E_{v_1} - U_{1(2)})\hat{I}_v & 0 \\
0 & 0 & 0 & 0 & 0 & (E_{v_2} - U_{1(2)})\hat{I}_v
\end{pmatrix}. \quad (5)
\]

Here \(\hat{I}_v\) is the identity operator in the 2 \(\times\) 2 space of atomic \(p_x, p_y\) orbital components of \(v_1\) and \(v_2\).
$E_{c_1}, E_v, E_{v_1}$ and $E_{v_2}$ are the energy differences between the lowest conduction band and the $c_1, v, v_1$ and $v_2$ bands, respectively, (see Fig. 4). In addition, we take

$$
\delta \hat{H}_{11(22)} = \begin{pmatrix}
0 & 0 & 0 & 0 \\
0 & 0 & 0 & 0 \\
-i b_{54} (k \cdot \Lambda)^T & 0 & 0 & 0 \\
-i \lambda_{46} (s \times \Lambda)^\dagger & -i b_{16} (k \cdot \Lambda)^T & 0 & 0
\end{pmatrix},
$$

Here $1 \times 2$ matrices $\Lambda_x = [0,1]$ and $\Lambda_y = [1,0]$ operate in the $p_x, p_y$ orbital components of $v_1$ and $v_2$ valence bands and the coefficients $b_{45}$, $b_{16}$ and $b_{15}^{cv}$ characterise the $c - v_1$, $v - v_2$ and $c_1 - v_2$ intra-layer couplings (associated with interband optical transitions excited by the in-plane polarised photons). Spin Pauli matrices $s_{x,y}$ produce spin flips upon the interband mixing which can be rooted into account linear in momentum interband terms in the monolayer Hamiltonian, discussed earlier in relation to the optical selection rules for the interband transitions

$$
\hat{H}_{12}^{(0)} = \begin{pmatrix}
t_{cc}^\Gamma & 0 & 0 & 0 \\
0 & 0 & 0 & 0 \\
0 & t_{cv}^\Gamma & 0 & 0 \\
0 & 0 & 0 & 0
\end{pmatrix},
$$

$$
\delta \hat{H}_{12} = \begin{pmatrix}
0 & (t_{cc_1} + \delta_{c_1c}) & (t_{cv}^\Gamma + \delta_{cv}) & 0 & 0 \\
(-t_{cc_1} + \delta_{c_1c}) & 0 & 0 & 0 & 0 \\
(-t_{cv}^\Gamma + \delta_{cv}) & 0 & 0 & 0 & 0 \\
0 & 0 & 0 & 0 & 0
\end{pmatrix}.
$$

The first of them describes the resonant interlayer hybridization of separately lower conduction and the top valence band edges, which was identified as the strongest hybridization effect, determined by the substantial weight of $s$ and $p_z$ chalcogen orbitals in the sub-lattice composition of the band edge states. The second term takes into account interband interlayer hybridization, which produces a much weaker effect on the band edge states. The second term takes into account interband interlayer hybridization, which produces a much weaker effect on the band edge states.

According to the table in Fig. 4, the on-layer states in bands $c$ are odd under $z \rightarrow -z$ reflection while bands $v$ and $c_1$ are even under the same transformation. Because of this, for a mirror symmetric arrangement of the layers, the corresponding interband interlayer couplings would obey the relation $t_{cv}^\Gamma = -t_{cv}^\Gamma$ and $t_{c_1c} = -t_{cc_1}$. To capture the mirror plane symmetry breaking for $\gamma$-stacking, we introduce parameters $\delta_{\alpha \beta}$ such that $t_{cc_v}^\Gamma = t_{cv}^\Gamma + \delta_{cv}, t_{cc_v}^\Gamma = -t_{cv}^\Gamma + \delta_{cv}, t_{cv_1} = t_{cv_2} = t_{cv}, t_{cv_1} = -t_{cv_2} + \delta_{cv_1v_2}, t_{c_1c_1} = -t_{c_1c_2} + \delta_{c_1c_2}$, and $t_{c_1c_2} = -t_{c_1c_2} + \delta_{c_1c_2}$. Overall, the $z \rightarrow -z$ symmetry breaking in the bilayer (which gives rise to the 2D SOC in the lowest conduction subband of the bilayer) is produced by the interplay between $\delta H_{11}$ and the contributions from $\delta_{\alpha \beta}$ in Eq. (9). For this we use $3^{rd}$ order perturbation theory with respect to parameters $\delta_{c_1c}, \delta_{cv}, b_{54}, b_{16}, b_{15}^{cv}, \gamma_{15},$ and $\gamma_{46}$, and this results in the spin-orbit coupling constant,

$$
\alpha_0 = 2 \left( \frac{b_{54} \lambda_{15} \delta_{cv}}{\Delta E_{cv_1} \Delta E_{g_1}} + \frac{b_{16} \lambda_{46} \delta_{cv}}{\Delta E_{cv_2} \Delta E_{g_1}} + \frac{b_{15}^{cv} \lambda_{46} \delta_{c_1c}}{\Delta E_{c_1c} \Delta E_{cv_2}} \right).
$$

## Notes

1. $\Lambda_x = [0,1]$ and $\Lambda_y = [1,0]$ operate in the $p_x, p_y$ orbital components of $v_1$ and $v_2$ valence bands.
2. The coefficients $b_{45}$, $b_{16}$ and $b_{15}^{cv}$ characterise the $c - v_1$, $v - v_2$ and $c_1 - v_2$ hybridizations.
3. Spin Pauli matrices $s_{x,y}$ produce spin flips upon the interband mixing.
4. Linear in momentum interband terms in the monolayer Hamiltonian are discussed earlier in relation to the optical selection rules.
5. The on-layer states in bands $c$ are odd under $z \rightarrow -z$ reflection while bands $v$ and $c_1$ are even under the same transformation.
6. Mirror symmetric arrangement of the layers results in the following interlayer couplings:
   - $t_{cc_v}^\Gamma = t_{cv}^\Gamma + \delta_{cv}$
   - $t_{cc_v}^\Gamma = -t_{cv}^\Gamma + \delta_{cv}$
   - $t_{cv_1} = t_{cv_2} = t_{cv}$
   - $t_{cv_1} = -t_{cv_2} + \delta_{cv_1v_2}$
   - $t_{c_1c_1} = -t_{c_1c_2} + \delta_{c_1c_2}$
   - $t_{c_1c_2} = -t_{c_1c_2} + \delta_{c_1c_2}$
7. The 2D SOC in the lowest conduction subband of the bilayer is produced by the interplay between $\delta H_{11}$ and the contributions from $\delta_{\alpha \beta}$ in Eq. (9).
8. $\alpha_0$ is the spin-orbit coupling constant defined by Eq. (9).
Here we also account for asymmetry induced by an external electric field so its effect on the on-layer energy of the orbitals in Eq. (5), captured by $\Delta E_{cv} \equiv -(t_{cc}^v + E_{cv})$, $\Delta E_{cv2} \equiv -(t_{cc}^v + E_{cv2})$ and $\Delta E_{cc1} \equiv -(t_{cc}^v + E_{cc1})$ are the energy differences between the lowest conduction subband and $v_1$, $v_2$ and $c_1$ bands while $\Delta E_{cc1} = -(t_{cc}^v + E_{cc1} + E_{cv}) = t_{cv} + E_{cv}$ is the energy difference between the lowest conduction subband and the 1st or 2nd topmost valence subband, respectively.

In the absence of external electric field, $U_1 = U_2 = 0$, and using parameters in Table III, we estimate that $\varepsilon_1(\alpha_0 = 0) = 0.35 \text{Vnm}^{-1}$. The dependence on a perpendicularly applied electric field $\varepsilon_2$ is approximated by

$$\varepsilon_2 = \left. \frac{d\alpha}{d\varepsilon_x} \right|_{U_1=U_2=0} = \left( \frac{\lambda_1^v + \lambda_4^v}{E_{cc1}} \right) \frac{2\varepsilon_2}{\Delta E_{cc1}} \left( \frac{1}{2\varepsilon_2} - \frac{1}{2\varepsilon_2} \right).$$

Here, $\Delta E_{c1(2)}$ and $\Delta E_{v1(2)}$ are the energy shifts of the $c$ and the $v$ bands in the 1st and 2nd layer respectively; $\Delta E_{v1(2)}$ and $\Delta E_{v2(1)}$ are the energy shifts of the bands $v_1$ and $v_2$ and $I^\nu$ is the identity operator in the $2 \times 2$ space of atomic $p_\alpha, p_\beta$ orbital components of the $c_1$ and $c_2$ bands. The terms $\Upsilon_{cv}$ and $\Upsilon_{v_1v_2}$ are responsible for $c-v$ and $v_1-v_2$ band mixing in the top layer: the interfacial $z \to -z$ symmetry breaking couples states of opposite parities. In the bottom surface, the interfacial effect is inverted, which is the reason for the inverted signs, $-\Upsilon_{cv}$ and $-\Upsilon_{v_1v_2}$ of the corresponding terms in $\delta \mathbf{H}_{11(22)}^{(1)}$. In Table II, we quote values of all those parameters obtained using DFT modelling described in Section II. In order to extract those parameters, the wavefunctions of bands $c$ and $v_1$ were obtained for the three different atomic arrangements described in Section II. By comparing their wavefunction distribution with the DFT-computed wavefunctions of suspended monolayer InSe, the mixing terms between opposite $z$-parity bands $\Upsilon_{cv}$ and $\Upsilon_{v_1v_2}$ was extracted for each configuration. Finally, from the DFT energy eigenvalues, the shifts in energy of bands $c$ and $v$ were obtained for each of the three different configurations; the energy shifts of bands $v_1$ and $v_2$ were neglected due to the very weak interlayer hybridization of those bands which results in a negligible contribution to the conduction band SOC strength. Using perturbation theory, we calculate the contribution of these additional terms towards bilayer SOC and find that the dominant effect comes from the $c - v$ band mixing, resulting in,

$$\alpha^{(1)} = \left[ \frac{b_{45} \lambda_1^v}{\Delta E_{cv} \Delta E_{cv2}} + \frac{b_{16} \lambda_4^v}{\Delta E_{cv} \Delta E_{cv2}} \right] \left( \Upsilon_{cv} - \Upsilon_{cv}^b \right).$$

The above equation suggests that encapsulation of InSe with the same material in the top and bottom would
result in the cancellation of the main part of such an additional contribution. Due to misalignment or an offset of the encapsulating crystals, this cancellation would never be exact leaving a residual effect due to the variation of InSe and, e.g., hBN stacking. Taking into account the random nature of such a variation, in the mechanically assembled structures, we estimate characteristic size of the residual SOC contribution using the characteristic difference of the Υ-valence parameters for two InSe/hBN stackings analyzed in Section II (Configuration 1 and 2 in Table II and Fig. 3). This gives $|\alpha^{(f)}| \sim 3.5 \text{meVÅ}$, which is an order of magnitude smaller than $\alpha_{0} = 13\text{meVÅ}$. As a result, for InSe bilayer encapsulated with hBN on both sides, the value and displacement field dependence of SOC can be well described using Eq. (9) and (10).

IV. SPIN-ORBIT COUPLING IN MULTILAYER InSe

Here, we combine the analysis of two factors that determine the strength of SOC in multilayer γ-InSe: the asymmetry embedded into the interlayer hybridization and the effect of an externally controllable electric field.

A. Self-consistent analysis of subband electrostatics in doped multilayer InSe films

In this section, the effect of an externally applied electrostatic potential (gating) for electrons in the lowest conduction subband is calculated self-consistently, and its effect on the charge distribution and on the band gap is discussed for the dual and single-gated FET geometry as sketched in insets of Fig. 11 and Fig. 13, respectively. To quantify the SOC in the lowest conduction subband of few-layer InSe films, we describe the subband structure of the latter (both dispersion and wavefunctions) taking into account the electrostatic potential profile induced by doping and gating. Our ‘workhorse’ is a 2-band hybrid k·p tight-binding (HkpTB) model previously discussed in Ref. 35, formulated in the basis of conduction, c and valence, v band states in each layer $[c_{1}, v_{1}, c_{2}, v_{2}, ...]$. The HkpTB Hamiltonian has the form,

$$
H_{ kpTB}^{N} \approx \begin{pmatrix}
\frac{\hbar^{2}k^{2}}{2m_{c}} + U_{1} & 0 & t_{cc} & t_{cv} & \cdots \\
0 & E_{v} + U_{1} & -t_{cv} & t_{vv} & \cdots \\
t_{cc} & -t_{cv} & \frac{\hbar^{2}k^{2}}{2m_{c}} + U_{2} & 0 & \cdots \\
t_{cv} & t_{vv} & 0 & E_{v} + U_{2} & \cdots \\
0 & 0 & t_{cc} & 0 & \cdots \\
\vdots & \vdots & \vdots & \vdots & \ddots
\end{pmatrix}
$$

(13)

Here, $t_{cc(vv)}$ parameterize the interlayer conduction-conduction (valence-valence) hops ($t_{cc} \equiv t_{c1c1}^{cc} + t_{c2c2}^{cc}$), while $t_{cv}$ ($t_{cv} \equiv t_{c1v1}^{cv} + t_{c2v2}^{cv}$) is the conduction to valence band hop. The zero of energy is set to the monolayer conduction band edge, so that $E_{c} \approx -2.8 \text{eV}$ is the energy of the monolayer’s topmost valence band at the Γ-point. We neglect the valence band dispersion in InSe monolayers, as earlier studies have shown that it is approximately flat over a large central part of the Brillouin zone. We also neglect any k-dependence in $t_{cv}$ for the same reason. The terms $U_{\eta}$ account for the electrostatic potential in layer $\eta$, and they are calculated as:

$$
U_{\eta>1} = U_{1} + e\alpha_{z} \sum_{\kappa=2}^{N_{\eta}} E_{(\kappa-1)\kappa},
$$

(14)

where $\alpha_{z} = 8.32 \text{Å}$ is the distance between adjacent layers and $E_{(\kappa-1)\kappa}$ is the electric field between layers $\kappa - 1$ and $\kappa$. $E_{(\kappa-1)\kappa}$ is obtained from the electron density on each InSe layer, $n_{\eta}$, as

$$
E_{(\kappa-1)\kappa} = \frac{e}{\varepsilon_{z} \varepsilon_{0}} \sum_{\eta=0}^{N} n_{\eta},
$$

(15)

where $N$ is the total number of InSe layers in the device, $n_{\eta}$ is the carrier concentration at the $\eta^{th}$ layer and $\varepsilon_{z}$ is the dielectric constant of InSe in the z-direction. We then approximate the electric field across a single layer as the mean of the fields either side of it,

$$
E_{\kappa} \simeq (E_{(\kappa-1)\kappa} + E_{(\kappa+1)\kappa})/2.
$$

(16)

Values of the parameters in the above Hamiltonian are listed in Table III. They are obtained by fitting the results of the numerical analysis of the 14-band model described in Ref. 33, 35, 55, and 56. It is also common, in order to obtain more flexibility in gating, to have both a back gate and a top gate applied to the device as shown in the dual-gated geometry in the inset of Fig. 11. To demonstrate the behaviour of the SOC coefficient in the dual-gated case, we reproduce the gating configuration used for transport experiments on a six-layer device studied in Ref. 11. In that work, a fixed positive top gate voltage was applied to dope the system. At $V_{bg} = 0$, the carrier density in the InSe films was measured to be $n_{c} \sim 4 \times 10^{12} \text{cm}^{-2}$ indicating that the charge density in the top plate was that same amount. To include a fixed top gate in our electrostatic calculations, we amend Eq. (15) to read

$$
E_{(\kappa-1)\kappa} = \frac{e}{\varepsilon_{z} \varepsilon_{0}} \left[ \sum_{\eta=0}^{N} n_{\eta} - n_{tg} \right],
$$

(17)

where $n_{tg}$ is the fixed top gate carrier density and $n_{\eta}$ the carrier density in layer $\eta$. In considering the single-gated
TABLE III. Dependence of the energy gap and of the effective mass of the lowest conduction subband as a function of the number of layers \( L \); \( m_0 \) is the free electron mass.

| \( L \) | Band gap (eV) | \( m_e/m_0 \) |
|---|---|---|
| 1 | 2.87 | 0.266 |
| 2 | 2.14 | 0.220 |
| 3 | 1.83 | 0.204 |
| 4 | 1.67 | 0.197 |
| 5 | 1.58 | 0.192 |
| 6 | 1.52 | 0.189 |
| 7 | 1.48 | 0.187 |
| 8 | 1.46 | 0.186 |
| 9 | 1.44 | 0.185 |
| 10 | 1.42 | 0.184 |

First, there are the intralayer dipole moments which mix wavefunctions of opposite parities within each layer under an applied electric field. Second, there is an interplay between the intrinsic inversion asymmetry of the lattice structure of \( \gamma \)-InSe, and the overall wavefunction \( z \to -z \) symmetry breaking due to the applied electrostatic potential. For the analysis of SOC in multilayer InSe it is necessary to include deeper valence bands \( v_1 \) and \( v_2 \) dominated by the \( p_z, p_y \) orbitals necessary for atomic SOC mixing with the \( p_z \) orbitals in \( c \) and \( v \) (see the orbital composition of each band in the character table on top of Fig. 4). On including the deeper valence bands, the hybrid \( \mathbf{k} \cdot \mathbf{p} \) tight-binding Hamiltonian \( \hat{H} \) of an \( N \)-layer InSe in the vicinity of the \( \Gamma \)-point \((k_x, k_y \to 0)\) previously discussed in Section III is rewritten as the sum of an unperturbed \( \hat{H}^{(0)} \) and a perturbative part \( \delta \hat{H} \),

\[
\hat{H} = \hat{H}^{(0)} + \delta \hat{H}. \tag{18}
\]

Writing the wavefunction eigenstates of the multilayer Hamiltonian \( \hat{H} \) in a 14 \( \times \) \( N \) band basis as \( \Psi = [\Phi_1, \Phi_2, \Phi_3, \Phi_4, ..., \Phi_N] \), where \( \Phi_w \) is the 14-band monolayer basis in layer \( w \) defined as

\[
\Phi_w = \begin{bmatrix} c_1^w, c_2^w, c_1^w, c_2^w, v_1^w, v_2^w, v_1^w, v_2^w, v_1^w, v_2^w, v_1^w, v_2^w \end{bmatrix},
\]

yields the following expression for \( \hat{H}, \hat{H}_0 \) and \( \delta \hat{H} \).
Here, indices $\eta$ and $\kappa$ label layers. The basis of each matrix $\hat{H}_{\eta\kappa}^{(0)}$ and $\hat{\delta H}_{\eta\kappa}$ is the 14-band monolayer InSe basis.

In $\hat{H}_{\eta\eta}^{(0)}$, $U_{\eta}$ is the electrostatic potential in the $\eta^l$ layer, $E_v$ is the monolayer topmost valence band energy as previously defined in the 2-band model, $E_v$, $E_u$ and $E_{c_1}$ are the energies of the $v_1 \ v_2$ and $c_1$ bands, and $\hat{\mathbf{I}}_\nu$ is the identity operator in the space of atomic $p_x \ p_y$ orbitals.

In $\hat{H}_{(\eta-1)\eta}^{(0)}$, parameters $t_{ccv}^\Gamma$ and $t_{cvv}^\Gamma$ are the neighbouring conduction-conduction (valence-valence) interlayer hoppings; no spin index has been included in Eq. (20d) and in Eq. (20b) as all non-zero matrix elements are spin independent. In $\hat{\delta H}_{(\eta-1)\eta}$, $t_{cvv}^\Gamma$ and $\delta_{ccv}$ are the $z$-symmetric and $z$-antisymmetric $c-v$ mixing interlayer hoppings, respectively (see Appendix A). In $\delta H_{\eta\eta}$, $d_{cv}$, $d_{vivv}$ and $d_{ccv}$ are the out-of-plane dipole moments (see Fig. 4). Coefficients $b_d$, $b_{16}$ and $b_{16}^{cv}$ are $\mathbf{k} \cdot \mathbf{l}$ mixing terms between $c - v_1$, $v - v_2$ and $c_1 - v_2$ respectively, while $\lambda_{16}$ and $\lambda_{15}$ are the atomic orbital SOC strengths for $c - v_2$ and $v - v_1$ spin-flip mixing, with values given in Table I. The latter is included using spin matrices $\mathbf{s}_x$ and $\mathbf{s}_y$. Matrices $\mathbf{A}_y$ and $\mathbf{A}_x$ are $1 \times 2$ matrices $[0, 1]$ and $[1, 0]$, respectively, operating in the $p_x \ p_y$ orbital component of the $v_1$ and $v_2$ valence bands and $\mathbf{k} \cdot \mathbf{A} \equiv k_x \mathbf{A}_x + k_y \mathbf{A}_y$ and $s \times \mathbf{A} \equiv s_x \mathbf{A}_x - s_y \mathbf{A}_x$.

In the absence of interband hoppings, and having neglected the interlayer hoppings between the deeper valence bands $v_1$ and $v_2$ and between band $c$ and the upper conduction band $c_1$, the subband eigenstates formed by $H_0$ define the orthogonal basis used in the Löwdin projection. The eigenstates of the $j^l$ conduction and valence subband states in this unperturbed Hamiltonian therefore have the form $|c\rangle = \sum_{\eta=1}^{\eta=N} \alpha_\eta^j |c_\eta\rangle$, $|v\rangle = \sum_{\eta=1}^{\eta=N} \beta_\eta^j |v_\eta\rangle$, $|v^{(2)}\rangle = |v_1^{(2)}\rangle$, $|c_1\rangle = |c_{1\eta}\rangle$ where $|c_\eta\rangle$, $|v_{\eta}\rangle$, $|v^{(2)}\rangle$ and $|c_1\rangle$ are the $c$, $v$, $v^{(2)}$ and $c_1$ monolayer eigenstates in layer $\eta$, respectively. In the following analysis we will only focus on the lowest conduction subband $\alpha_\eta^1 \equiv \alpha_\eta$. For the purpose of calculating the SOC coefficient as a function of carrier density, the $v_1$, $v_2$ and $c_1$ subbands are approximated as all being located at $E^{c}_{v_1} = E_{v_1} - U_{av}$, $E^{c}_{v_2} = E_{v_2} - U_{av}$ and $E^{c}_{c_1} = E_{c_1} - U_{av}$ respectively, where $U_{av}$ is the average electrostatic potential per layer. This is due to a small change in the on site electrostatic potential, $\Delta U_{\eta-1} = U_{\eta-1} - U_{\eta}$, as compared with the $E_{v_1}$, $E_{v_2}$ and $E_{c_1}$ energy denomina-
Defining, the perturbation theory have to be third-order in the expansion. Consequently, the lowest-order non-zero terms in the perturbation theory have the form, 

\[ \Delta H_{m_m'}^{(3)} = -\frac{1}{2} \sum_{l,m',m''} H_{m_m} H_{m'm} H_{m''m'} \left( E_{m'} - E_l \right) \left( E_{m''} - E_l \right) \]

\[ - \frac{1}{2} \sum_{l,m',m''} H_{m_m} H_{m'm} H_{m''m'} \left( E_{m'} - E_l \right) \left( E_{m''} - E_l \right) \]

\[ + \frac{1}{2} \sum_{l,l'} \frac{H_{m_m} H_{l'l'} H_{l'm'}}{E_m - E_l} \left( E_m - E_{l'} \right) \]

\[ + \frac{1}{2} \sum_{l,l'} \frac{H_{m_m} H_{l'l'} H_{l'm'}}{E_m - E_l} \left( E_m - E_{l'} \right), \]

(22)

where the \( m, m' \) indices correspond to \( \hat{H}_0 \) subband eigenstates in block A and the \( l, l' \) index to any subband eigenstate in block B (see Appendix D). Energies \( E_m(l) \) correspond to the energy of the \( m^{th} \) or \( l^{th} \) eigenstate. Contributions to SOC originate from the 3-step loop Feynman diagrams in Fig. 7-10, with spin reversed initial and final states \( c \uparrow \langle \downarrow \rangle \)

The Feynman diagrams, originating from the inversion asymmetric parameter \( \delta_{cv} \) and \( \delta_{c}\hat{c}v \) in combination with the mixing with deeper valence bands and SOC as shown in Fig. 7 and Fig. 8, give a term,

\[ \Delta H_{11}'' = 2 \left[ \sum_{j=N}^{j=1} \sum_{e=N}^{e=1} \left( \frac{b_{44} \lambda_{15} \delta_{cv}}{\Delta E_{c\uparrow 1} \Delta E_{e\downarrow 1}} + \frac{b_{16} \lambda_{46} \delta_{cv}}{\Delta E_{c\downarrow 2} \Delta E_{e\uparrow 2}} \right) \right] \times \alpha_{\xi} \left( \beta_{\xi+1}^{2} + \beta_{\xi-1}^{2} \right) \sum_{\eta=1}^{\eta=N} \alpha_{\eta} \beta_{\eta}^{2} + \sum_{\eta=1}^{\eta=N} \left( \frac{b_{46} \lambda_{46} \delta_{c\hat{c}v}}{\Delta E_{c\uparrow 1} \Delta E_{c\downarrow 1}} \right) \times \alpha_{\eta} \alpha_{\eta} \left( \alpha_{\eta+1} + \alpha_{\eta-1} \right) \left( s \times k \right), \]

(23)

where \( \delta_{cv} \) and \( \delta_{c}\hat{c}v \) is the \( z \)-asymmetric parameters between \( c \) and \( v \) and between \( c_1 \) and \( c \) defined in Eq. (20d) and further discussed in Appendix A. In the presence of an external electrostatic potential, the signs of \( \delta_{cv} \) and \( \delta_{c}\hat{c}v \) become important, as it can be related to placing a single electrostatic gate on one of the surfaces and the orientation (up/down) of externally controlled electric field, \( E_z \).

The two diagrams in Fig. 9, give a SOC term in the

![Diagram](Image) FIG. 7. (Left) Feynman diagram of the interlayer spin-flip loops due to the \( \gamma \)-stacking involving the upper conduction band \( c_1 \). (Right) Feynman diagram of the interlayer spin-flip loops due to the \( \gamma \)-stacking involving the deeper valence bands \( v_1 \) and \( v_2 \). Such contribution is only relevant for the Dresselhaus SOC in the valence band \( v \) as shown in Appendix A. Doted lines (\( \cdots \)) label the terms in \( \delta H \) responsible for inversion symmetry breaking. Dashed lines (\( \cdots \)) label the intra-atomic SOC mixing between different bands. Solid lines label the \( \mathbf{k} \cdot \mathbf{p} \) interband mixing terms in \( \delta H \). Different colors label pairs of loops that produce competing contributions in the same order of perturbation theory.

![Diagram](Image) FIG. 8. Feynman diagram of the interlayer spin-flip loops due to the \( \gamma \)-stacking responsible for breaking the \( z \rightarrow -z \) symmetry in the \( c \) to \( v \) hopping parameters \( t_{cv} \) and \( t_{vc} \). Doted, dashed and solid lines follow the same convention as in Fig. 7.
form of,
\[
\Delta H'_1 = 2 \left[ \sum_{j=1}^{N} \sum_{k=1}^{N} \left( \frac{b_{13} \lambda_{15} t_{1r}^{1r}}{\Delta E_{cv1} \Delta E_{cvj}^{1r}} + \frac{b_{16} \lambda_{46} t_{1r}^{1r}}{\Delta E_{cv1} \Delta E_{cvj}^{1r}} \right) \times \alpha_{k} (\beta_{j+1}^{l} - \beta_{j-1}^{l}) \right] (s \times k),
\]
where \(\alpha_{k}\) and \(\beta_{j}^{l}\) are the components of the lowest conduction subband and the \(j^{th}\) valence subband respectively; \(k\) labels the layer index. \(\Delta E_{cvj}^{1r} \equiv E_{c} - E_{v}^{j}\) is the energy difference between the lowest conduction subband and the \(j^{th}\) valence subband and \(\Delta E_{cvj}^{1(2)} \equiv E_{c} - E_{v}^{j(2)}\) is the energy gap between the lowest conduction subband and the \(v_1\) and \(v_2\) subbands located at \(E_{v1(2)} - U_{av}\). The loops shown in Fig. 10 for the dipolar mixing terms give a SOC term in the form of
\[
\Delta H''_{1} = 2 \left[ \sum_{j=1}^{N} \sum_{k=1}^{N} \left( \frac{d_{cv \lambda_{15} b_{54}}}{\Delta E_{cv1} \Delta E_{cvj}^{1r}} + \frac{d_{cv \lambda_{46} b_{16}}}{\Delta E_{cv1} \Delta E_{cvj}^{1r}} \right) \times (s \times k),
\]
where \(d_{cv}\) is the matrix element of the out-of-plane dipole operator between the monolayer conduction and valence bands, \(d_{v1v2}\) is the out-of-plane dipole moment between \(v_1\) and \(v_2\) and \(d_{c1c}\) the out-of-plane dipole between \(c_1\) and \(c\). \(E_{\eta}\) is defined as the electric field in layer \(\eta\) and \(\Delta E_{cc1} \equiv E_{c} - E_{c1}\) is the energy difference between the lowest conduction subband and the set of \(c_1\) subbands located at \(E_{c1} - U_{av}\). In accounting for the dipolar terms, some care must be taken in choosing its sign in the few-layer case, as is further explained in Appendix B.

Combining all these contributions enables us to describe the dependence of SOC strength, \(\alpha\), on the number of layers, electric field, and doping in the film as shown in Figs. 11, 12, and 13. For example, as illustrated in Fig. 12, in multilayer InSe in a single-gated FET, doping the device to carrier densities \(> 10^{13} \text{cm}^{-2}\) can lead to the compensation of the intrinsic SOC by the contribution of the gate-induced electric field.

C. SOC analysis in InSe films using a quantum well model

To describe thicker films, it is more practical to use a quantum well model for InSe films. For this, we describe the dispersion of electrons in the \(\mathbf{k} \cdot \mathbf{p}\) theory expansion near the A-point conduction band edge of bulk InSe as
\[
E_{c}(p_{z}) = \left( \frac{\hbar^2}{2m_{A}} + \frac{\xi p_{z}^2 a_{p}^2}{2m_{A}} \right) p_{z}^2 + \frac{\hbar^2 p_{z}^2}{2m_{A}} + \alpha_{\infty} \left( 1 - \frac{\chi a_{e}^2 p_{z}^2}{\pi^2} \right) (s \times \mathbf{p}),
\]
where \(m_{A}\) and \(m_{Ae}\) are the in-plane and out-of-plane effective mass at the A-point and the parameters \(\xi\) and \(\chi\) take into account the anisotropic non-parabolicity of the electron’s dispersion characteristic for layered systems.
FIG. 11. SOC strength dependence on displacement field and carrier density for a six-layer InSe dual-gated FET device as shown in the inset. The solid and dotted lines indicate respectively when the applied displacement field suppresses or enhances the Dresselhaus SOC.

In Fig. 13(b) we show the $p_z$-dependence (around the A-point) of the linear in $k_x, k_y$ spin-orbit coupling computed by DFT for bulk InSe using QSGW approach, to compare with the SOC form in Eq. (26). This has to be complemented with the generalised Dirichlet-Neumann boundary conditions for the quantum well wavefunction $\Psi(z)$ at the encapsulating interfaces,

$$\Psi \pm \nu a_z \partial_z \Psi = 0, \quad \nu \approx 1.42.$$  \hspace{1cm} (27)

The latter determines the values for the wave numbers of the electron’s standing waves,

$$p_z = \frac{n\pi}{(N+2\nu)a_z};$$  \hspace{1cm} (28)

which determines the subband and layer-number-dependence of the subband mass and SOC parameter,

$$\frac{1}{m_n} \approx \frac{1}{m_A} \left(1 - \frac{6.2n^2}{(N+2\nu)^2}\right),$$  \hspace{1cm} (29)

$$\alpha_{n|N}(n << N) \approx \alpha_\infty \left(1 - \frac{\chi}{(N+2\nu)^2}\right).$$

By fitting $\alpha_{1|N}$ described in Eq. (2) to the values of the lowest subbands SOC strength in Fig. 13 we find that $\alpha_\infty = 34.5$ meVÅ and $\chi = 14.9$ respectively. Additionally, the results of the calculations, performed in the same films subjected to an electric field $E_z$ perpendicular to the layers shown in Fig. 1 and 11 show an approximately linear SOC strength dependence on $E_z$. We describe the latter as

$$\alpha(N, E_z) = \alpha(N) - \beta(N) E_z,$$  \hspace{1cm} (30)

with the values of $\beta(N)$ for $N \geq 2$ shown in the inset of Fig. 1. Further to the DFT calculations for the few-layer case, in Fig. 3(b) we use previous quasiparticle self-consistent GW (QSGW) calculations for bulk $\gamma$-InSe to extract the $k_z$-dependence of the coefficient of the linear component of SOC for small in-plane momentum near $k_x=k_y=0$, for both the conduction and valence bands. This shows that as $k_z$ approaches the bulk band edge (located at $k_z = \pi/a_z$) the SOC strength increases, implying that as $k_z$ is restricted by confinement in thin films of InSe, the SOC strength can be expected to decrease from its bulk value, with smaller strengths for thinner films.

V. MAGNETOTRANSPORT STUDIES OF InSe FILMS IN THE FET GEOMETRY AND THEIR COMPARISON WITH THEORY

In order to probe the nature of SOC in InSe, we fabricated a dual-gated multiterminal 6 layer $\gamma$-InSe device using mechanical exfoliation and hexagonal boron nitride (hBN) encapsulation, which were carried out in an inert atmosphere of a glovebox. Such encapsulation was
FIG. 13. (a) SOC coefficient $\alpha$ computed for InSe films with various thicknesses and carrier densities in a single-gated FET geometry (inset), calculated assuming $\epsilon_z = 9.9$ for the InSe. The data shown in circles ($\bigcirc$) were obtained by exact diagonalization of the 14-band Hamiltonian in Ref. 55 and compared with the perturbation theory results obtained by Löwdin partitioning ($\bigtriangleup$) (note that for $N=1$, $\alpha = 0$ for $n_e = 0 \times 10^{12}$ cm$^{-2}$). Inset shows the usual configuration of a single-gated FET device. The solid and dashed lines indicate the fitted dependence of the Dresselhaus term as a function of the number of layers when the quantum well approximation holds ($N \geq 4$) and when it does not respectively. (b) In red, the QSGW-calculated SOC strength as a function of $p_z$ in bulk $\gamma$-InSe for the conduction band$^{56,59}$. As $p_z$ approaches the band edge located at $p_z = \frac{\pi}{a_z} = 0.378$ Å$^{-1}$, the SOC strength increases following a quadratic dependence on momentum $p_z$ shown in green ($\alpha(p_z) = \alpha_\infty \left( 1 - \frac{\chi a_z^2 p_z^2}{2} \right)$, where $\alpha_\infty = 34.5$meVÅ and $\chi = 14.9$). This increasing trend indicates that for greater confinement under a decreasing number of layers, a weaker linear Dresselhaus SOC is expected.

FIG. 14. (a) Optical photograph of an encapsulated InSe flake (light blue) equipped with few-layer graphene (FLG) contacts (red). Yellow polygons illustrate gold leads connecting FLG. Green colour corresponds to the bottom hBN flake deposited on top of an oxidized Si wafer (dark blue). (b) Calculated charge density distribution along the different layers in the dual-gated device under study. At a carrier concentration of $n_e = 8 \times 10^{12}$ cm$^{-2}$, the distribution of charges becomes $z \rightarrow -z$ symmetric as the top plate carrier density is fixed at $n_{tg} = 4 \times 10^{12}$ cm$^{-2}$.
FIG. 15. Weak antilocalization feature in conductivity measured in a 6-layer dual-gated InSe device with corresponding optimal fits (Black). Carrier densities were measured in the range from $1.7 \times 10^{12} \text{ cm}^{-2}$ to $0.1 \times 10^{12} \text{ cm}^{-2}$. Blue indicates the upper and lower bound fits of the corrections to magnetoconductivity. Top inset shows the carrier density at each back gate voltage obtained from Hall-effect measurements. The finite carrier density at $V_{bg}=0$ is due to the applied top gate voltage corresponding to $V_{tg}=8 \text{ V}$. The linear relation between the carrier density and the back gate voltage for a fixed top gate of $V_{tg}=8 \text{ V}$ was found to be $n_e = \Upsilon (V_{bg} - V'_{bg}(V_{tg}=8 \text{ V}))$ where $V'_{bg}(V_{tg}=8 \text{ V}) = -67.6 \text{ V}$ and $\Upsilon = 5.71 \times 10^{10} \text{ V}^{-1} \text{ cm}^2$.

Using Drude formula we determined the mean free path of charge carriers, $\lambda$, and respective scattering time, $\tau$, important parameters critical for further analysis. The effective mass for the lowest conduction subband used to extract $\tau$ was $m_c = 0.12 m_e$, obtained from an accurate calculation of the bulk effective mass accounting both for electron-electron and electron-phonon interaction effects in the bulk conduction band.\(^{63}\)

An experimental manifestation of the SOC strength can be found in the weak antilocalization (WAL) corrections to magnetoconductance\(^{64-67}\) produced by the interference of electron waves propagating along closed loops of random walks\(^{68,69}\). Such behavior has been observed in recent studies of few-layer single-gated GaSe\(^{70}\) and InSe\(^{71,72}\).

In Ref. 72, the fitting procedure used to extract the SOC strength from the corrections to magnetoconductance was the formalism developed by Hikami, Larkin and Nagaoka\(^{66}\) for systems where the spin relaxation mechanism is dominated by scattering with magnetic impurities\(^{73,74}\). As the $\gamma$-stacked phase in InSe is non-centrosymmetric and therefore the spin relaxation mechanism is expected to be Dyakonov-Perel, their extracted spin relaxation parameters from WAL fits were overestimated. In Ref. 71 the enhancement of the SOC as compared to our estimated bulk SOC strength value at the band edge ($\alpha_\infty \approx 34 \text{ meVÅ}$) is a result of an impurity deposition layer formed at the interface of the suspended device; this forms a sharp potential barrier at the interface and therefore increases the SOC strength.

From our weak antilocalization measurements, the spin and phase relaxation times can be obtained by fitting the corrections to conductivity with respect to these two parameters in the range of magnetic fields where the minimum in magnetoconductance appears.

The WAL corrections to the conductivity of the 6-layer device with the same characteristics as reported in Ref. 11 were measured as a function of the magnetic field with 1 mT magnetic field step. As shown in Fig. 15, at mag-
nentic fields 10–30 mT, a clear minimum in the magnetoconductance is observed. The corrections to conductivity $\Delta \sigma(B) - \Delta \sigma(0)$ were measured in the range of 0–90 mT, and both the spin and phase relaxation time were fitted with the formalism developed by Iordanskii, Larkin and Pitaevskii\textsuperscript{75,76} (ILP) for systems where the lack of inversion symmetry leads to the electron’s spin precessing and to relaxation by Dyakonov-Perel mechanism. Such formalism was used for carrier densities $< 2 \times 10^{12}$ cm$^{-2}$; above that carrier densities, the assumption of the ILP formalism was used for carrier densities and to relaxation by Dyakonov-Perel mechanism. Such Pitaevskii with the formalism developed by Iordanskii, Larkin and

\[ \sigma_{\text{back}} = -\frac{e^2}{2\pi^2\hbar} \left( \frac{\lambda}{l_B} \right)^2 \sum_{N'=0}^{\infty} \left( T_{\hat{A}_{N'}} (I - \hat{A}_{N'})^{-1} - \frac{P_{N'}}{1 - P_{N'}} \right), \] (31)

\[
\hat{A}_{N'} \equiv \left[ \begin{array}{cc}
P_{N'-2} - S_{N' - 2}^{(0)} & R_{N' - 2}^{(1)} \\
S_{N' - 2}^{(2)} & P_{N' - 1} - 2S_{N' - 1}^{(0)} & R_{N' - 1}^{(1)} \\
S_{N' - 2}^{(2)} & R_{N' - 1}^{(1)} & P_{N'} - S_{N'}^{(0)}
\end{array} \right],
\]

\[
P_{N'} \equiv \frac{l_B}{\lambda} \int_0^\infty \exp \left( -\frac{l_B}{\lambda} x - \frac{x^2}{2} \right) L_{N'}(x^2) dx,
\]

\[
S_{N'}^{(0)} \equiv \frac{l_B}{\sqrt{2\lambda}} \sqrt{\frac{N'!}{(N' + \mu)!}} \int_0^\infty \exp \left( -\frac{l_B}{\lambda} x - \frac{x^2}{2} \right) x^\mu L_{N'}^\mu(x^2) \sin^2 \left( \frac{\Omega \tau l_B}{\lambda} \right) dx,
\]

\[
R_{N'}^{(1)} \equiv \frac{l_B}{\sqrt{2\lambda}} \sqrt{\frac{N'!}{(N' + \mu)!}} \int_0^\infty \exp \left( -\frac{l_B}{\lambda} x - \frac{x^2}{2} \right) x^\mu L_{N'}^\mu(x^2) \sin \left( 2\Omega \tau l_B \right) dx,
\]

Here, $l_B \equiv \sqrt{\frac{\hbar}{eB}}$ is the magnetic length, and in Eq. (31), $\hat{I}$ is defined as $\hat{I} \equiv \frac{\lambda}{\tau + \tau_{\phi}}$ where $\tau_{\phi}$ is the phase relaxation time. The precession frequency is related to the spin relaxation time $\tau_{SO}$ through $\frac{1}{\tau_{SO}} = 2l^2 r$. As done previously with the ILP formalism, both the phase and spin relaxation times were taken as fitting parameters. In Fig. 16, the inverse proportionality between the spin relaxation time and the diffusion coefficient $D$ confirms that the spin relaxation mechanism is Dyakonov-Perel\textsuperscript{79,80}. From $\tau_{SO}$, the SOC coefficient is extracted and compared with our theoretical calculation in Fig. 18. In Fig. 18 the SOC coefficient at different carrier densities was calculated at the experimentally established dielectric constant $\varepsilon_z = 9.9$ for InSe\textsuperscript{92}. Very good agreement was found between the calculated SOC coefficient and the experimentally extracted SOC strength. Furthermore, by looking at the two different branches originated from the orientation of the crystal being parallel or antiparallel to the applied electric field, it was found that at a carrier density of $n_c = 8 \times 10^{12}$ cm$^{-2}$ the two branches converged at a single point. This indicates no dependence neither on crystal orientation nor on electrostatic profile.

In Fig. 17, the parallel orientation $z\rightarrow z$, the magnetoconductance obtained using the few-layer HkpTB study and a quantum well model give the matching results, and the theoretically computed SOC strengths are compared with the results of weak antilocalization measurements on dual-gated multilayer InSe films showing a good agreement between theory and experiment.

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VI. CONCLUSION

Overall, the description of SOC strength (as a function of the number of layers and the applied electric field piercing the multilayer film) obtained using the few-layer HkpTB study and a quantum well model give the matching results, and the theoretically computed SOC strengths are compared with the results of weak antilocalization measurements on dual-gated multilayer InSe films showing a good agreement between theory and experiment.

The size of SOC constant we compute for InSe films with 2-10 layers thickness is comparable to the SOC strength in quantum wells of conventional semiconduc-
FIG. 17. (a-d) SOC strength as a function of the displacement field and carrier density for different number of layers. Crystal orientation is chosen such that the applied displacement field counteracts the Dresselhaus SOC when the displacement field is positive. The dark black lines indicate the disappearance of SOC due to the application of a displacement field which compensates the SOC from the intrinsic lack of inversion symmetry in the different multilayers.

FIG. 18. SOC coefficient $\alpha$ as experimentally extracted from weak antilocalization measurements of the dual-gated 6 layer device\textsuperscript{11}, compared to the value obtained in the self-consistent calculation. The blue dashed line indicates the value of $\alpha$ in the absence of any electrostatic gating and doping. The same notation for the solid and dotted lines is used as in Fig. 1,12 and 11.

tors, such as GaAs, InAs, HgTe. What makes 2D InSe different from those spintronic systems is that the SOC strength in it can be tuned over a wide range. Additionally contribution originating from the asymmetry of an hBN/InSe interface was analysed and shown to be negligible (as compared with the intrinsic SOC in the film) for InSe encapsulated in hBN both on top and in the bottom, and also to decay as $N^{-3}$. Moreover we demonstrate that spin-orbit coupling strength for electrons near the conduction band edge in few-layer $\gamma$-InSe films can be tuned over a wide range, from $\alpha=0$ to $\alpha \approx 70$ meVÅ. This tunability illustrated in Fig. 17 for the films of various thicknesses is the result of a competition between film-thickness-dependent intrinsic and electric-field-induced SOC, potentially, allowing for electrically switchable spintronic devices. As shown in Fig. 17 and Fig. 1, displacement fields in the range of 1-2 Vnm\textsuperscript{-1} can turn the SOC on and off.

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monolayer conduction and valence under $\sigma_z$ when combined with the opposite lattices of selenium atoms on the outside of each layer. This causes the hops to be inversion symmetric, which when combined with the opposite $z$-symmetries of the monolayer conduction and valence under $\sigma_z$ reflection (i.e. $z \rightarrow -z$ symmetry) gives $t_{cv} = -t_{vc}$, $t_{c1c} = -t_{c1c}$, and $t_{v1v2} = -t_{v1v2}$. It is transparent from Eq. (1) that inversion symmetry would prohibit the existence of extrinsic SOC. Consequently, we require terms in our model which break inversion symmetry (such as an applied electric field or the interlayer pseudopotentials arising from the $\gamma$-stacking). The indium atoms provide such an asymmetry-in-the $\gamma$ stacking there is a vertically opposite interlayer In/Se pair heading in one direction along the $z$-direction, while in the other direction the indium atom is opposite an empty space in the adjacent layer. In the $k \cdot p$ model, the effect of this symmetry breaking is to give $t_{cv}$, $t_{c1c}$ and $t_{v1v2}$ slightly different magnitudes as compared with $-t_{cv}$, $-t_{c1c}$ and $-t_{v1v2}$, so we define three new parameters: $2\delta_{cv} \equiv t_{cv} + t_{vc}$, $2\delta_{c1c} \equiv t_{c1c} + t_{c1c}$, and $2\delta_{v1v2} \equiv t_{v1v2} + t_{v1v2}$. In order to obtain the parameters $\delta_{cv}$ and $\delta_{c1c}$ relevant for the analysis of the Dresselhaus SOC in the conduction band, the linear SOC splittings at each individual $k_z$ are obtained from the QUESTAAL package by linearly fitting the energy differences between the two spin split bands (see Fig. 19). Firstly, the parameters $\delta_{cv}$ and $\delta_{c1c}$ were fitted for the $\alpha$ vs $k_z$-dependence of band $c$ (red curve in Fig. 19), and then the $\delta_{v1v2}$ parameter was fitted from the $\alpha$ vs $k_z$-dependence of band $v$ (green curve in Fig. 19). Using the same perturbative analysis as in Section IV in the bulk limit, the Dresselhaus SOC at each $k_z$ is obtained both for the $c$ and $v$ bands respectively, namely

$$\alpha_c(p_z) = 4 \cos(p_z a_z) \left( \frac{\delta_{cv} b_{54} \lambda_{15}}{(E_c - E_v)(E_c - E_{v1})} + \frac{\delta_{cv} b_{16} \lambda_{46}}{(E_c - E_v)(E_c - E_{v2})} \right) + \left( \frac{\delta_{c1c} b_{54} \lambda_{15}}{(E_c - E_v)(E_c - E_{v1})} + \frac{\delta_{c1c} b_{16} \lambda_{46}}{(E_c - E_v)(E_c - E_{v2})} \right).$$

(A1)

and

$$\alpha_v(p_z) = 4 \cos(p_z a_z) \left( \frac{\delta_{cv} b_{54} \lambda_{15}}{(E_v - E_c)(E_v - E_{v1})} + \frac{\delta_{cv} b_{16} \lambda_{46}}{(E_v - E_c)(E_v - E_{v2})} \right).$$

where $p_z = \vec{p} \cdot -k_z$. The fitting parameters considered are the terms $\delta_{cv}$, $\delta_{c1c}$, $\delta_{v1v2}$ and $\lambda_{46}$ as the 14-band fit applied to the InSe bulk dispersion did not account for any them. The optimal parameters found in order to fit the spin splitting vs $k_z$-dependence in the vicinity of the band edge where perturbation theory is best applicable were $\delta_{cv} = 0.014$ eV, $\delta_{c1c} = 0.022$ eV, $\delta_{v1v2} = -0.001$ eV and $\lambda_{46} = -0.09$ eV.

![FIG. 19. (Solid) Bulk SOC as a function of $p_z$ for $c$ and $v$ bands. (Dots) SOC strength at different $p_z$ obtained from the perturbative analysis in Eq. (A1) and (A2).](image)

Appendix B: Determination of the signs of $d_{cv}$, $d_{v1v2}$ and $d_{c1c}$

While on their own the signs of $d_{cv}$ and $t_{cv}$ may be chosen arbitrarily through an appropriate choice of basis in the monolayer Hamiltonian, the product of $d_{cv}$ and $t_{cv}$ does not have such degree of freedom. In order to determine the relative signs of the different dipole moments, it is necessary to look at their $k$-dependence as we move away from the $\Gamma$-point. In considering the conduction to valence band interlayer hopping (both the $z$-symmetric and $z$-antisymmetric) as a perturbation to our conduction or valence subband wavefunctions, the $k$-dependence of the bilayer valence band dipole moment follows easily.

Appendix A: Determination of parameters $\delta_{cv}$, $\delta_{c1c}$ and $\delta_{v1v2}$ from bulk SOC

For the calculation of subband energies and dispersions, it was sufficient to approximate the interlayer hops as being entirely between the inversion symmetric sublattices of selenium atoms on the outside of each layer. The optimal parameters found in order to fit $\alpha$ vs $k_z$-dependence in the vicinity of the band edge where perturbation theory is best applicable were $\delta_{cv} = 0.014$ eV, $\delta_{c1c} = 0.022$ eV, $\delta_{v1v2} = -0.001$ eV and $\lambda_{46} = -0.09$ eV.

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as:

\[ \langle v_{2L}|ez|v_{2L} \rangle = 2\delta_{cv} \left[ \frac{d_{cv}}{E_{g2L}} + \frac{t_{cv}e_0}{2E_{g2L}E'_{g2L}} \right] \]  

for the valence band, and

\[ \langle c_{2L}|ez|c_{2L} \rangle = -2\delta_{cv} \left[ \frac{d_{cv}}{E_{g2L}} + \frac{t_{cv}e_0}{2E_{g2L}E'_{g2L}} \right] \]  

for the bilayer conduction band. In the above equation, \( d_{cv} = ||c|ez|v|| = 1.68 \text{ eÅ} \) is the matrix element of the out-of-plane dipole operator between the monolayer conduction and valence bands and \( v_{2L} \) and \( c_{2L} \) are the topmost valence subband and lowest conduction subband wavefunctions in a bilayer system at the Γ-point. \( E'_{g2L} = E_c - E_v - (t_{cc} - t_{vv}), \ E''_{g2L} = E_c - E_v + t_{cc} + t_{vv}, \) and \( E''_{g2L} = E_c - E_v - (t_{cc} + t_{vv}) \) are the energy differences between the bilayer bands in the absence of the interband hoppings. Comparison of these expressions with the signs of the quantities calculated using DFT gives, for a choice of positive \( t_{cv} \) and negative \( d_{cv} \), a positive \( \delta_{cv} \) when the \( +z \) direction is chosen such that the vertical In-Se interlayer pair in the interface between two layers the Se atom lies above the In atom in the \( \gamma \)-stacking. Conversely, a negative \( \delta_{cv} \) is obtained for the opposite orientation. On calculating perturbatively the value of \( d_{cv} \) at a finite \( k \), the following result is obtained

\[ d_{cv}(k) \equiv \langle v|ez|c \rangle = \langle v_0|ez|v_0 \rangle + \frac{b_{54}b_{16}k^2d_{v1v2}}{\Delta E_{v1v2}\Delta E_{v2v}}. \]  

By looking at the negative trend of \( |d_{cv}| \) and the hybrid \( k \cdot p \) tight-binding values quoted in Table III, it is transparent that if \( d_{cv} \) is positive \( d_{v1v2} \) is as well positive. Furthermore, if \( d_{cv} \) is negative, the value of \( d_{v1v2} \) should be negative as well. In order to find the sign of the dipole moment \( d_{c1c} \) a similar perturbative analysis is applied for \( d_{v1v2} \):

\[ d_{v1v2}(k) \equiv \langle v_1|ez|v_2 \rangle = \langle v_{1,0}|ez|v_{2,0} \rangle + \frac{b_{54}b_{16}k^2d_{cv}}{\Delta E_{v1c}\Delta E_{v2v}} \]

\[ + \frac{b_{54}b_{16}k^2d_{c1c}}{\Delta E_{v1c}\Delta E_{v2v}}. \]

In comparing the red and the green curve in Fig. 20, the much more pronounced steepness of the red curve as compared to the green curve at low values of \( k \) indicates that \( d_{c1c} \) must be negative for a positive \( d_{v1v2} \) and vice versa.

**Appendix C: Interfacial contribution to multilayer InSe SOC**

In addition to the crystalline and the electrostatically induced \( z \rightarrow -z \) asymmetry, few-layer InSe is a material sensitive to interfacial effects due to its limited thickness. Such effects may have an impact in the SOC strength of multilayer InSe and must therefore be taken into consideration. The same two InSe-hBN configurations used for the analysis of interfacial effects in bilayer InSe shown in Table II (configuration 1 and 2) were also used for the calculation of the interface-induced SOC in multilayer InSe as their contribution in the absence of an external electrostatic potential is only dependent on the encapsulating substrates and on the film thickness. Interface effects are taken into account by adding into the multilayer Hamiltonian two additional contributions identical to Eq. (11). Firstly, bands \( c \) and \( v \) with a relevant Se \( p_z \) orbital composition, experience in the outer layers a shift in energy due to the interaction with the \( p_z \) orbitals of the encapsulating hBN. Therefore, an additional energy shift is added to the \( c,v,v_1 \) and \( v_2 \) bands of the 1st and the \( N \)th layer.

Additionally, the hBN interfaces break \( z \rightarrow -z \) symmetry in the outer layers mixing bands with opposite \( z \)-parity but identical in-plane symmetries. The following perturbative term accounting for these effects is introduced in the multilayer Hamiltonian.

\[ \delta \hat{H}_{11(NN)}^{(I)} = \]  

\[ \begin{pmatrix} 
\Delta E_{c1(N)} & 0 & \pm \Upsilon^{t/b}_{cv} & 0 & 0 \\
0 & \Delta E_{v1(N)} & 0 & 0 & 0 \\
\pm \Upsilon^{t/b}_{cv} & 0 & 0 & 0 & 0 \\
0 & 0 & 0 & \Delta E_{v1(N)} & \hat{t}_v \\
0 & 0 & \pm \Upsilon^{t/b}_{v1v2} & \hat{t}_v & \Delta E_{v21(N)} 
\end{pmatrix}, \]

where \( \Upsilon^{t/b}_{cv} \) and \( \Upsilon^{t/b}_{v1v2} \) are the mixing terms between bands \( c \rightarrow v \) and \( v_1 \rightarrow v_2 \) in the top interface and \( -\Upsilon^{t/b}_{cv}, -\Upsilon^{t/b}_{v1v2} \) in the bottom interface.
are ones mixing bands $c - v$ and $v_1 - v_2$ at the bottom interface. Note that such mixing terms require an opposite sign due to the opposite sign due to the opposite direction of the interfacial effective electric fields at the two InSe/hBN interfaces. Given the very small interfacial energy shift of bands $c$ and $v$ and the very weak hybridization between bands $v_1$ and $v_2$, the dominant contribution to the conduction band SOC strength originates from the interfacial terms mixing bands of opposite parity (see Fig. 22). Among them, the most relevant contribution originates from the term $\Upsilon_{cv}$ mixing bands $c$ and $v$, which, in the absence of an applied electric field, yields to the following contribution to the SOC strength,

$$\Delta H^{(l)}_{11} = 2\left(\frac{b_{15}\lambda_{15}}{\Delta E_g^{1}\Delta E_{cv}^{1}} + \frac{b_{16}\lambda_{16}}{\Delta E_g^{2}\Delta E_{cv}^{2}}\right)(\Upsilon_{cv}^{l}x_{1}\beta_{1} - \Upsilon_{cv}^{l}x_{N}\beta_{N}) (s \times k),$$

where $\Delta E_g^{1}$ is the energy between the lowest conduction subband and the topmost valence band (i.e. the energy gap) and $\Delta E_{cv}^{l}$ is the energy difference between the lowest conduction subband and the $v_1(2)$ subbands. The number of layers dependence of the interfacial SOC strength can be extracted expanding $\Delta E_g^{1}$, $\Delta E_{cv}^{1,2}$, $\alpha_{1(2)}$ and $\beta_{1(2)}$ as a function of the number of layers in the quantum well approximation presented in Ref. 35. In such approximate framework, the out-of-plane wavevector $k_z$ depends on the number of layers as $k_z = \frac{\pi}{a_z} + \frac{n\pi}{(N+2\nu)a_z}$ and the wavefunctions for both the conduction and the valence bands are approximated as the eigenstates of a quantum well size $L = (N + 2\nu)a_z$ ($\Phi_{c}^{n} \approx \Phi_{v}^{n} \approx \sqrt{\frac{1}{(N+2\nu)a_z}} \cos\left(\frac{n\pi}{(N+2\nu)a_z}\right)$). From this quantum well model, a $\frac{1}{(N+2\nu)}$ dependence of the interfacial SOC strength is expected, as confirmed by the fit presented in Fig. 21. Given the smallness of the interfacial SOC strength compared to the layer-number dependent Dresselhaus SOC, any contribution coming from the hBN/InSe interface will be neglected for the rest of our analysis.

**Appendix D: Löwdin partitioning method**

In order to obtain the 3rd order corrections to the hybrid $k \cdot p$ tight-binding Hamiltonian, the standard method of Löwdin partitioning is applied. The total multilayer Hamiltonian is written in the basis of the unperturbed subbands eigenstates obtained from diagonalizing the $\hat{H}_0$ part of the Hamiltonian in Eq. (20b),

$$\hat{H} = \hat{H}_0 + \hat{H}'$$

where $\hat{H}'$ is the perturbative part, namely the projection of $\delta \hat{H}$ in the orthogonal subband basis formed by $\hat{H}_0$ ($H_{\rho\omega}' \equiv \langle \rho | \delta \hat{H} | \omega \rangle$). In the partitioning method, two diagonal blocks are defined, A and B and a unitary transformation is applied to the entire Hamiltonian matrix in order to remove the non-block-diagonal elements. The set $A$ is defined as the elements within the lowest con-
for the SOC splitting shown in Figs. 7-9 and 10, the

elements within B. Having identified the loops responsible
where (1 and 2 refer to the

\[ H = H_0 + H_1 + H_2. \]  

(D5)

Transforming the Hamiltonian with a unitary transformation of the form \( e^S \),

\[ \hat{H} = e^{-S} \hat{H} e^S, \]  

(D6)

duction subband \( e^1 \)

\[ \hat{H}_A = \begin{pmatrix} \langle c^1| \hat{H}_A |c^1 \rangle & \langle c^1| \hat{H}_A |c^2 \rangle \\ \langle c^1| \hat{H}_A |c^2 \rangle & \langle c^1| \hat{H}_A |c^1 \rangle \end{pmatrix}, \]  

(D2)

while the set B are the matrix elements within the valence
subbands or the upper conduction subbands,

\[ \hat{H}_B = \begin{pmatrix} \langle v^1| \hat{H}_B |v^1 \rangle & \langle v^1| \hat{H}_B |v^2 \rangle & \cdots \\ \langle v^2| \hat{H}_B |v^1 \rangle & \langle v^2| \hat{H}_B |v^2 \rangle & \cdots \\ \vdots & \vdots & \ddots \end{pmatrix}, \]  

(D3)

where the numerical indices such as 1 and 2 refer to the
1st or 2nd subbands. The non-block-diagonal elements, \( H_{nbd} \) are the elements mixing the terms of the A and B block namely

\[ \hat{H}_{nbd} = \begin{pmatrix} \langle c^1| \hat{H} |v^1 \rangle & \langle c^1| \hat{H} |v^2 \rangle & \cdots \\ \langle v^1| \hat{H} |c^1 \rangle & \langle v^1| \hat{H} |c^2 \rangle & \cdots \\ \langle v^2| \hat{H} |c^1 \rangle & \langle v^2| \hat{H} |c^2 \rangle & \cdots \end{pmatrix}. \]  

(D4)

The expression in Eq. (D1) is rewritten in terms of \( \hat{H}_1' \) (the matrix containing the perturbations within block A and B), and \( \hat{H}_2' \) (the non-zero perturbations between sets A and B)

\[ \hat{H} = \hat{H}_0 + \hat{H}_1' + \hat{H}_2'. \]  

The deeper valence band states are projected into the lowest conduction subband. From the definition of the A block, the matrix elements \( \langle c^1| \hat{H} |c^1 \rangle \) and \( \langle c^1| \hat{H} |c^1 \rangle \) are the terms responsible for the SOC splitting. We get the following expressions for the block and non-block-diagonal matrix elements,

\[ \hat{H}_{bd} = \sum_{j=0}^{\infty} \frac{1}{(2j)!}[\hat{H}^{(0)} + \hat{H}^{(1)}, \hat{S}]^{(2j)} + \sum_{j=0}^{\infty} \frac{1}{(2j+1)!}[\hat{H}^{(2)}, \hat{S}]^{(2j+1)}, \]

\[ \hat{H}_{nbd} = \sum_{j=0}^{\infty} \frac{1}{(2j+1)!}[\hat{H}^{(0)} + \hat{H}^{(1)}, \hat{S}]^{(2j+1)} + \sum_{j=0}^{\infty} \frac{1}{(2j)!}[\hat{H}^{(2)}, \hat{S}]^{(2j)}. \]  

(D7)

The non-block-diagonal terms are then set to 0 forcing the third order in the perturbation Hamiltonian (\( \Delta H^{(3)} \)) to be

\[ \Delta H_{mm'}^{(3)} = -\frac{1}{2} \sum_{l,m'} \left[ \frac{H_{ml}^H H_{l'm'}^H H_{l'm}^H}{(E_{m'} - E_l)(E_{m'} - E_l)} + \frac{H_{ml}^H H_{l'm'}^H H_{l'm}^H}{(E_{m} - E_l)(E_{m'} - E_l)} \right] \]  

(D8)

The non-block-diagonal terms are then set to 0 forcing the third order in the perturbation Hamiltonian (\( \Delta H^{(3)} \)) to be

where \( (m, m') \) are elements within A and \( (l, l') \) are elements within B. Having identified the loops responsible for the SOC splitting shown in Figs. 7-9 and 10, the

mixing between the conduction and the deeper valence bands projected into the lowest conduction subband has the form,
\[
\Delta H_{11}^{(3)} = 2 \sum_{j=1}^{\eta=N} \sum_{\eta=1}^{N} \frac{\langle c^\dagger | \delta H | v_{2,\eta}^\dagger \rangle \langle v_{2,\eta}^\dagger | \delta H | v_{1,\eta}^\dagger \rangle \langle v_{1,\eta}^\dagger | \delta H | c^\dagger \rangle}{\Delta E_{cv1} \Delta E_{cv2}} + 2 \sum_{j=1}^{\eta=N} \sum_{\eta=1}^{N} \frac{\langle c^\dagger | \delta H | v_{1,\eta}^\dagger \rangle \langle v_{1,\eta}^\dagger | \delta H | v_{2,\eta}^\dagger \rangle \langle v_{2,\eta}^\dagger | \delta H | c^\dagger \rangle}{\Delta E_{cv1} \Delta E_{cv2}} 
\]

Knowing the origin of the 3-step loop processes described in Section IV, the Hamiltonian that contributes to the SOC in the absence of a relevant interfacial term can be decomposed as

\[
\Delta H_{11}^{(3)} = \Delta H_1' + \Delta H_1'' + \Delta H_1''', \tag{D10}
\]

where the different terms correspond to the different mechanisms behind SOC in band \(c\),

\[
\Delta H_1' = 2 \left[ \sum_{j=1}^{\eta=N} \sum_{k=1}^{N} \left( \frac{b_{54} \lambda_{15} c_{15}^\dagger}{\Delta E_{cv1} \Delta E_{cv2}} + \frac{b_{16} \lambda_{46} c_{46}^\dagger}{\Delta E_{cv1} \Delta E_{cv2}} \right) \alpha_k (\beta_{k+1} - \beta_k) \left( \sum_{\xi=1}^{\eta=N} \alpha_{\xi} \beta_{\xi} \right) (s \times k), \tag{D11}
\]

\[
\Delta H_1'' = 2 \left[ \sum_{j=1}^{\eta=N} \sum_{k=1}^{N} \left( \frac{\mathcal{E}_v d_{cv1} \lambda_{15} b_{54}}{\Delta E_{cv1} \Delta E_{cv2}} + \frac{\mathcal{E}_v d_{cv1} \lambda_{46} b_{16}}{\Delta E_{cv1} \Delta E_{cv2}} \right) (\alpha_k \beta_k) \left( \sum_{\xi=1}^{\eta=N} \alpha_{\xi} \beta_{\xi} \right) + \sum_{\eta=1}^{\eta=N} \left( \frac{b_{16} \lambda_{46}}{\Delta E_{cv1} \Delta E_{cv2}} \right) (s \times k), \right.
\]

\[
\Delta H_1''' = 2 \left[ \sum_{j=1}^{\eta=N} \sum_{k=1}^{N} \left( \frac{b_{54} \lambda_{15} \delta_{cv}}{\Delta E_{cv1} \Delta E_{cv2}} + \frac{b_{16} \lambda_{46} \delta_{cv}}{\Delta E_{cv1} \Delta E_{cv2}} \right) \alpha_k (\beta_{k+1} + \beta_k) \left( \sum_{\xi=1}^{\eta=N} \alpha_{\xi} \beta_{\xi} \right) + \sum_{\eta=1}^{\eta=N} \left( \frac{b_{16} \lambda_{5} \delta_{cv1}}{\Delta E_{cv1} \Delta E_{cv2}} \right) (s \times k), \right.
\]

Finally, using Eq. (D9) the interfacial contribution to the SOC strength coming from the dominant \(\gamma_{cv}^{l/b}\) term in Eq. (C2) has the form

\[
\Delta H_{11}^{(l)} = 2 \left[ \sum_{j=1}^{\eta=N} \left( \frac{b_{54} \lambda_{15}}{\Delta E_{cv1} \Delta E_{cv2}} + \frac{b_{16} \lambda_{46}}{\Delta E_{cv1} \Delta E_{cv2}} \right) \left( \gamma_{cv}^{l} \alpha_{l} \beta_{l} - \gamma_{cv}^{b} \alpha_{N} \beta_{N} \right) \left( \sum_{\xi=1}^{\eta=N} \alpha_{\xi} \beta_{\xi} \right) \right) (s \times k), \tag{D12}
\]

Considering the limit where the applied electric field is zero, this term simplifies to

\[
\Delta H_{11}^{(l)} = -2 \left[ \frac{b_{54} \lambda_{15}}{\Delta E_{cv1} \Delta E_{cv2}} + \frac{b_{16} \lambda_{46}}{\Delta E_{cv1} \Delta E_{cv2}} \right] \left( \gamma_{cv}^{l} \alpha_{l} \beta_{l} - \gamma_{cv}^{b} \alpha_{N} \beta_{N} \right) (s \times k). \tag{D13}
\]
Note that the SOC coefficients, $\alpha$, implied by the DFT results, 18.6 meVÅ and 32.3 meVÅ for bilayer and trilayer respectively, are somewhat larger than those predicted by the model presented in this work - this is due to the substantial underestimation of the band gap of InSe by DFT. We therefore base the parametrization of the model on GW results for bulk crystals.