Comparing the effective enhancement of local and non-local spin-orbit couplings on honeycomb lattices due to strong electronic correlations

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(Dated: November 9, 2021)

We investigate the interplay of electronic correlations and spin-orbit coupling (SOC) for a one-band and a two-band honeycomb lattice model. The main difference between the two models concerning SOC is that in the one-band case the SOC is a purely non-local term in the basis of the \( p_z \) orbitals, whereas in the two-band case with \( p_x \) and \( p_y \) as basis functions it is purely local. In order to grasp the correlation effects on non-local spin-orbit coupling, we apply the TRILEX approach that allows to calculate non-local contributions to the self-energy approximately. For the two-band case we apply dynamical mean-field theory. In agreement with previous studies, we find that for all parameter values in our study, the effect of correlations on the spin-orbit coupling strength is that the bare effective SOC parameter is increased. However, this increase is much weaker in the non-local than in the local SOC case. Concerning the TRILEX method, we introduce the necessary formulas for calculations with broken SU(2) symmetry.

I. INTRODUCTION

The interplay of spin-orbit coupling (SOC) and strong electronic correlations has attracted a lot of attention in recent years [1].

On the one hand, mainly triggered by the discovery of insulating behavior in Sr\(_2\)IrO\(_4\) [2], a lot of research has been devoted to understand, how SOC can influence the degree of correlations in materials with heavy transition metal ions [3–7]. In these iridate compounds, for instance, SOC has been identified as the driving force to reach the strongly correlated insulating state [3, 8]. Without SOC, the relevant electronic orbitals would be the \( t_{2g} \) orbitals. The SOC now splits this manifold into \( J_{eff} = 3/2 \) and \( J_{eff} = 1/2 \) orbitals, where the latter ones become half-filled, leading to a strongly-correlated spin-orbit polarized electronic state [6].

On the other hand, one can also ask the inverse question, which is to what extent the spin-orbit coupling is changed by the inclusion of electronic correlations. This is also important in the context of correlated topological insulators [1]. In topological insulators in general, SOC is the main driving force for topological properties, and different topological phases can occur as function of the spin-orbit coupling parameter \( \lambda_{SO} \), interaction strength \( U \), and/or other parameters like crystal or external field effects [9]. If parameters are such that the system is close to a phase transition, external perturbations might be able to even switch between topological and non-topological states, as shown, e.g., in Ref. [10].

Quite generally, it has been found both in model as well as material-related studies [7, 11–16] that electronic correlations enhance the SOC strength, leading to an effective SOC \( \lambda_{eff} > \lambda_{SO} \). For ruthenates, this has also been confirmed experimentally [17]. We want to note that the concept of effective single-particle parameters is not at all limited to the case of SOC presented here. For instance, band-width (hopping) renormalization due to non-local interactions [18, 19] has been studied.

In this work, we investigate the effective SOC for two similar, but yet complementary systems. First, we want to study this behavior for a model, where the spin-orbit coupling is a non-local term in the Hamiltonian. We choose the Kane-Mele-Hubbard (KMH) model [20–22] as an example. It has been a popular model to investigate on the effects of correlations to a topological phase [9, 23–47], but has never been studied concerning the enhancement of spin-orbit coupling due to correlations.

Second, we construct a two-band honeycomb model motivated by bismuthene [48, 49], which can be viewed as the two-orbital variant of the graphene honeycomb lattice, but where the SOC term is again local. Comparing these two cases with similar structure and both at half-filling, we can quantify how non-local spin-orbit couplings respond to electronic correlations as opposed to the case of spin-orbit couplings. As we will explain in more detail below, this enhancing effect is calculated from the zero-energy limit of the self-energy.

While in order to investigate the enhancement of a local spin-orbit coupling dynamical mean-field theory (DMFT) [50] works very well and is, hence, our method of choice for bismuthene, this is no longer sufficient for the KMH model. Due to the local approximation of the self-energy in DMFT, its contribution to the non-local SOC is a-priori zero. In order to get a non-zero contribution we have to go beyond the local approximation of DMFT. Therefore, we adopt the TRILEX approach developed by Ayral and Parcollet [51, 52], which uses a local approximation of the electron-boson vertex and, hence, recovers a non-local contribution to the self-energy. Since

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FIG. 1. Sketch of the lattice structure of the two honeycomb lattices. Top: Single-band honeycomb lattice for the KMH model with non-local SOC. Bottom: Two-orbital lattice, relevant for bismuthene with a local SOC term. The bonds with strongest hopping term in the models are indicated by $t$, and the SOC is also indicated by arrows.

The spin-orbit coupling term in the Kane-Mele Hubbard model breaks SU(2)-symmetry, we need to develop the necessary TRILEX formulas for this case, which is possible within the original TRILEX formulation. However, the SU(2)-broken version of this original TRILEX approach can violate hermiticity for channel-off-diagonal components. As detailed in Sec IV we therefore employ a variant of the method coined TRILEX $\Lambda^2$ [53] which resolves this issue.

The generalization of the TRILEX formulas to SU(2)-broken is not only necessary in the present case (where we do paramagnetic calculations with SOC) but also for other calculations inside a spin-symmetry broken phase, such as an antiferromagnet.

II. MODELS

A. One-band model: KMH

The KMH model describes locally interacting electrons on the two-dimensional honeycomb lattice with nearest-neighbor hopping and spin-orbit coupling. The Hamiltonian reads

$$H = -t \sum_{\langle ij \rangle, \sigma} c^\dagger_{i\sigma} c_{j\sigma} + U \sum_i n_{i\uparrow} n_{i\downarrow}$$

$$+ i\lambda^{1B}_{SO} \sum_{\langle\langle ij\rangle\rangle, \sigma, \sigma'} \nu_{ij} c^\dagger_{i\sigma} \sigma \sigma' c_{j\sigma'},$$  

with the creation (annihilation) operator $c^\dagger_{i\sigma}$ ($c_{i\sigma}$) creating (annihilating) an electron with spin $\sigma$ at site $i$. The density operator is $n_{i\sigma} = c^\dagger_{i\sigma} c_{i\sigma}$, $\sigma^z$ is the Pauli matrix, and $\nu_{ij}$ gives +1 for every right turn and −1 for every left turn as illustrated in Fig. 2. The first term is a simple tight binding term, including just next-nearest neighbors. The second term is the well-known (purely local) Hubbard interaction. The last term includes just next-nearest neighbors and relates to the SOC in graphene. Here SOC is a purely non-local quantity, and its strength is given by $\lambda^{1B}_{SO}$, where we use the superscript 1B to refer to the 1-band model.

The non-interacting case ($U = 0$) can be solved exactly by a Fourier transformation of the Hamiltonian. Therefore we are fixing the unit vectors of the lattice to $a_1 = (3/2, \sqrt{3}/2)$ and $a_2 = (0, \sqrt{3})$. The honeycomb lattice has a natural bipartite structure, i.e., it consists of two interpenetrating triangular sub-lattices A and B (see also Fig. 2). As the SOC term breaks SU(2) symmetry, a full basis-set for $H(k)$ consists of four different species with a defined spin-direction. Taking

$$\Psi_{k}^{(S_z)} = (a_{k,\uparrow}, b_{k,\uparrow}, a_{k,\downarrow}, b_{k,\downarrow})$$

as a basis-set with $a_{k,\sigma}$ living on sub-lattice A and $b_{k,\sigma}$ on sub-lattice B, the Hamilton matrix reads

$$H_0^{(S_z)}(k) = \begin{pmatrix}
\gamma_k & -g_k & 0 & 0 \\
-g_k & \gamma_k & 0 & 0 \\
0 & 0 & -\gamma_k & -g_k \\
0 & 0 & -g_k^* & \gamma_k
\end{pmatrix},$$

with

$$g_k = te^{i\frac{2\pi}{\sqrt{3}}k_y} \left[ e^{i\frac{\pi}{2}k_y} + 2 \cos \left( \frac{\sqrt{3}k_y}{2} \right) \right]$$
\[ \gamma_k = 2\lambda_{SO}^{B} \left[ 2\sin\left(\frac{\sqrt{3}}{2}k_y\right) \cos\left(\frac{3}{2}k_x\right) - \sin\left(\sqrt{3}k_y\right) \right]. \]  

The two inequivalent high-symmetry points lie at \( \mathbf{K} = (2\pi/3, 2\pi/(3\sqrt{3})) \) and \( \mathbf{K'} = (2\pi/3, -2\pi/(3\sqrt{3})) \). At these high-symmetry points, in the absence of \( \lambda_{SO}^{B} \) we find Dirac cones. Finite spin-orbit coupling opens up a gap of size \( 6\sqrt{3}\lambda_{SO}^{B} \), resulting in a topological phase. A more detailed discussion of the KMH Model can be found, e.g., in [22, 27].

B. Two-band model: Bismuthene

Bismuth has five valence electrons and, therefore, forms a buckled hexagonal lattice with all the \( p \)-orbitals being near the Fermi edge. When passivating this sheet of bismuth atoms with either a suitable substrate or hydrogen, not only does the buckling vanish and the system becomes planar, but the \( p_z \)-orbitals are also shifted away from the Fermi energy. This happens because the excess valence electron binds with the passivating molecules, resulting in Dirac cones at the \( K \)-points for the \( p_z \)-orbitals. As a result, the system has to be described as an effective two-band model. The theoretical background and experimental verifications of substrate-passivated bismuthene have been presented by Reis et al. [48]. In this work, the passivation is performed by adding hydrogen atoms to the bismuthene sheet, as was done by Freitas et al. [49]. Due to the high atomic number of bismuth, spin-orbit coupling is large and opens a considerable gap at the Fermi level. Note that performing the passivation with hydrogen atoms instead of using a SiC substrate results in a direct gap instead of an indirect one.

As already mentioned, the important effect of the passivation of bismuthene is that it removes the \( p_z \) orbital from the Fermi level, and the relevant bands are instead coming from \( p_x \) and \( p_y \) orbitals. In this two-orbital basis, it has been shown that the SOC term has local matrix elements and can be written as a local operator \([48]\),

\[ H_{SO} = \frac{1}{2} \begin{pmatrix} 0 & i & 0 & 0 \\ -i & 0 & 0 & 0 \\ 0 & 0 & 0 & -i \\ 0 & 0 & i & 0 \end{pmatrix}. \]  

The non-interacting Hamiltonian for our two-band model is constructed as follows. We perform density-functional theory calculations for bismuthene without SOC included using the VASP code [54–58]. To obtain the atomic positions as input data, we used the knowledge that bismuthene has a hexagonal structure just like graphene and took the atomic distances from Freitas et al. [49]. We added a hydrogen atom to each of the bismuth atoms, for sublattice A in the positive and for sublattice B in the negative \( z \)-direction. We used a \( \Gamma \)-centered \( 27 \times 27 \times 1 \) \( k \)-point grid and a basis-set cutoff \( \text{ENCUT} = 200 \text{eV} \). We then construct a non-interacting Hamiltonian \( H_{SO}^{\text{noSOC}}(\mathbf{k}) \) by a maximally-localized Wannier90 projection [59] to the \( p_z \) and \( p_y \) orbitals. We provide the output of this projection [60] in a machine-readable format in the supplementary material of this publication. Other formats of the data are available from the authors upon request. The full non-interacting Hamiltonian is then given by

\[ H_{0}(\mathbf{k}) = H_{0}^{\text{noSOC}}(\mathbf{k}) + \lambda_{SO}^{B} H_{SO}, \]  

where we use \( \lambda_{SO}^{B} \) as SOC strength and the superscript \( 2B \) to refer to the two-band model.

In order to check whether a purely local SOC term is a good approximation, we also did DFT calculations including the SOC directly. We find that by choosing \( \lambda_{SO}^{B} = 0.435 \text{eV} \) in Eq. (6) the agreement between the band structures obtained in the two ways is excellent. Approximately the same value for the SOC was derived in the work of Reis et al. [48]. The full non-interacting Hamiltonian is then used as input for the DMFT calculations. As interaction part of the Hamiltonian we use a Kanamori-Hamiltonian of the form

\[ H_{\text{int}} = U \sum_{m} n_{m\uparrow} n_{m\downarrow} + U' \sum_{m \neq m'} n_{m\uparrow} n_{m'\downarrow} + (U' - J) \sum_{m < m', \sigma} n_{m\sigma} n_{m'\sigma} + J \sum_{m \neq m'} c_{m\uparrow}^{\dagger} c_{m'\downarrow} c_{m'\uparrow} c_{m\downarrow} + J \sum_{m \neq m'} c_{m\uparrow}^{\dagger} c_{m'\uparrow} c_{m'\downarrow} c_{m\downarrow}, \]  

where \( U \) is the Coulomb interaction and \( J \) denotes the Hund’s coupling, and \( U' \) is defined as \( U - 2J \) due to rotational symmetry. For the calculations a we choose a fixed ratio of \( J = 0.2U \).

The DMFT calculation has been performed with the TRIQS library [61] and the continuous-time quantum Monte Carlo impurity solver as implemented in the TRIQS/CTHYB application [62].

C. Making the models comparable

Since we are calculating the non-interacting band structure for the two-band model for bismuthene from DFT, we first normalize this band structure such that the largest matrix element in the tight-binding Hamiltonian is \( t = 1 \). This largest element is the nearest-neighbor orbital-diagonol hopping, as shown in Fig. 1. This allows us to use this matrix element \( t = 1 \) as unit of energy throughout the paper.

Next, we have to set the SOC strengths in the two models. In both cases, we choose this strength such that the ratio of the gap \( \Delta \) opened by SOC to the relevant
paramagnetic phase has the following form:

interacting models are shown in Fig. 3. 

SO
for a model without SOC, the contribution \( \Sigma \) from DFT for Bismuthene, which in the same units reads \( \lambda \). The dispersion relations for the non-interacting two-band model (bismuthene) is \( \Delta \). Note that for the two-band model we use the band-width of the two innermost bands for this comparison. This results in \( \lambda^{1B} = 0.14t \) and \( \lambda^{2B} = 0.21t \). Note that this value for the SOC in the two-band case is very close to the actual value determined from DFT for Bismuthene, which in the same units reads \( \lambda^{DFT} = 0.24t \). The dispersion relations for the non-interacting models are shown in Fig. 3.

III. SELF-ENERGY CONTRIBUTION TO THE EFFECTIVE SOC

A. One-band model and non-local SOC

The self-energy of the KMH-model (Eq. (1)) in the paramagnetic phase has the following form:

\[
\Sigma^{(Sz)} = \begin{pmatrix}
\Sigma^d + \Sigma^{SO} & \Sigma_{AB} & 0 & 0 \\
\Sigma_{BA} & \Sigma^d - \Sigma^{SO} & 0 & 0 \\
0 & 0 & \Sigma^d - \Sigma^{SO} & \Sigma_{AB} \\
0 & 0 & \Sigma_{BA} & \Sigma^d + \Sigma^{SO}
\end{pmatrix}
\] (8)

Here, \( k \) and \( i\omega_n \) indices have been dropped to improve the readability. On the diagonal we find an ordinary diagonal part \( \Sigma^d \) and another part \( \Sigma^{SO} \) that breaks SU(2) symmetry.

The motivation to use this nomenclature is the following. For a model without SOC, the contribution \( \Sigma^{SO} \) is obviously zero and one is left only with \( \Sigma^d \). When SOC is included, we additionally get a contribution that adopts the symmetry of \( \gamma_k \) (Eq. (4)) in the Hamiltonian (2). The symmetry of these terms in the Brillouin zone matches \( \gamma_k = -\gamma_{-k} \), and in spin- and sublattice space we obtain a sign structure corresponding to the \( \text{diag}(\gamma_k, -\gamma_k, -\gamma_k, \gamma_k) \):

part of Hamiltonian (2). Furthermore, this structure also implies that the local contribution of \( \Sigma^{SO} \), i.e. summed over all momenta, is zero, and does not produce any spin splitting on the impurity/atom.

All TRILEX self-consistent calculations are done using the block structure of Eq. (8). However, in the post-processing we need to disentangle \( \Sigma^{SO} \) from \( \Sigma^d \) in order to define an effective SOC. To do so, we can rotate the spin-subspace from the vertical (2) into the horizontal axis (1) using the transformation matrix

\[
T = e^{-i\pi/4\nu_p} \times \begin{pmatrix} 1 & 0 \\ 0 & 1 \end{pmatrix}
\] (9)

This leads to the following matrix form of the self-energy

\[
\Sigma^{(Sz)} = T\Sigma^{(Sz)}T^{-1} = \begin{pmatrix}
\Sigma^d & \Sigma_{AB} & \Sigma^{SO} & 0 \\
\Sigma_{BA} & \Sigma^d & 0 & -\Sigma^{SO} \\
0 & -\Sigma^{SO} & \Sigma_{BA} & \Sigma^d \\
0 & 0 & \Sigma_{BA} & \Sigma^d + \Sigma^{SO}
\end{pmatrix}
\] (10)

where the contribution due to SOC is now well separated on the off-diagonal.

From this self-energy \( \Sigma^{SO} \) we can define an effective spin-orbit coupling. Some care has to be taken because of the sign factor \( \nu_{ij} \) in the Hamiltonian, which gives a factor +1 or −1 depending on the direction of the bond, see Fig. 2. We can define the static enhancement of the SOC as

\[
\lambda_{eff}^{1B} = \lambda_{SO}^{1B} + \nu_{ij} \text{Im} \Sigma^{SO}(\mathbf{r} - \mathbf{r}') = a_{ij}, i\omega = i\omega_0
\] (11)

with \( a_{ij} \) being the vector of the honeycomb lattice along a next-nearest neighbor bond, where \( \nu_{ij} = \pm 1 \). From the lattice structure it is obvious that this connecting vector is one of the two primitive unit-cell vectors, and depending on the direction of the bond one needs to take +1 or −1 for \( \nu_{ij} \). We will show during the discussion of the results that this choice is reasonable.

B. Two-band model and local SOC

Since the SOC term Eq. (5) in the two-band model is a local term, it suffices to look at the local self-energy. Although being local in terms of lattice coordinates, the local Hamiltonian as well as the local self-energy acquire spin-off-diagonal terms. The eigenbasis of the local non-interacting Hamiltonian Eq. (6) is given by the two basis functions

\[
| p^1\rangle = \frac{1}{\sqrt{2}} (| p^+ \rangle \pm i | p^- \rangle) \\
| p^\dagger \rangle = \frac{1}{\sqrt{2}} (| p^\dagger^+ \rangle \pm i | p^\dagger^- \rangle)
\] (12)

where the eigenvalues are \( \pm \lambda_{SO}^{2B} / 2 \), respectively. In this basis, the self-energy is diagonal. We denote the matrix elements by \( \Sigma^{\pm}_r \) with \( r = \pm \), which fulfill the relations
The electron-boson vertex function by a local one. More precisely, in the Heisenberg decoupling, the interaction term and approximating the electron-boson coupling vertex $\Lambda$ is approximated by a local quantity in the Hedin equations, and $\alpha$ and $\beta$ denote sublattice indices.

$$\Sigma^\sigma_{\alpha} (k, i\omega) = \alpha \frac{\Lambda^\sigma_{\alpha} (k, i\omega)}{\alpha} + \Sigma^\sigma_{\alpha} (k, i\omega) \delta_{\alpha}$

$P^\eta_{\alpha} (q, i\Omega) = \frac{\Lambda^\eta_{\alpha} (q, i\Omega)}{\beta} + P^\eta_{\alpha} (i\Omega) \delta_{\alpha}$

FIG. 4. Lattice self-energy $\Sigma(k, i\omega)$ (upper panel) and polarization $P(q, i\Omega)$ (lower panel) within TRILEX: the electron-boson coupling vertex $\Lambda$ is replaced by a local one. Similar to what has been done in three-band systems [7], we can write the self-energy in this basis as

$$\Sigma = \frac{1}{2} (\Sigma^+ + \Sigma^-) \mathbb{I} + (\Sigma^0 - \Sigma^\uparrow) \mathbf{l} \cdot \mathbf{s},$$

where $\mathbb{I}$ is the identity and $\mathbf{l} \cdot \mathbf{s}$ is the SOC operator. From this relation an expression for an effective spin-orbit coupling can be defined as [7]

$$\lambda^{2B}_{\text{eff}} = \lambda^{2B}_{\text{SO}} + \left( \Re \Sigma^0 (i\omega_0) - \Re \Sigma^\uparrow (i\omega_0) \right).$$

**IV. TRILEX ($\Lambda^2$)**

The triply irreducible local expansion [51, 52] (TRILEX) is a diagrammatic extension of DMFT [63] that was developed by Ayral and Parcollet and consists of decoupling the interaction term and approximating the electron-boson vertex function by a local one. More precisely, in the Heisenberg decoupling, the interaction term is rewritten as

$$U_{n\uparrow} n_{\downarrow} = \frac{1}{2} \sum_{l=0,x,y,z} U^l_{\alpha} n^l_{\alpha} n^l_{\alpha},$$

with

$$n^l_{\alpha} = \sum_{c} c_{\alpha}^l \sigma_{\alpha c}^l c_{\alpha}^l,$$

where $\sigma^0 = 1$ and $\sigma^{x/y/z}$ are the Pauli matrices. $U^l$ denotes the bare interaction in the channel $l$. This decoupling, however, is not unique (“Fierz ambiguity”) [51, 64], and the ratio between the bare interactions $U^l$ in the different channels are determined by a parameter $\alpha$, the so-called Fierz parameter:

$$U^{ch} = U(3\alpha - 1),$$

$$U^{x} = U^{y} = U^{z} = U(\alpha - \frac{2}{3}).$$

Here we have defined the bare interaction in the charge-channel as $U^{ch} \equiv U^0$. As the SOC term in the KMH model breaks SU(2) symmetry, we can assume spin-rotational invariance only in the xy-plane, leaving us with three independent channels:

$$ch = 0, x/y, z = z$$

Through Hubbard-Stratonovich transformations in each of the channels the electron-electron interaction problem is then transformed into an electron-boson coupling problem. The bosonic and fermionic lattice Green’s functions are given by the Dyson equations

$$G^\sigma (k, i\omega) = \left[ G^\sigma_0 (k, i\omega) \right]^{-1} - \Sigma^\sigma (k, i\omega)^{-1},$$

$$W^{\eta\eta'} (q, i\Omega) = U^\eta [\delta_{\eta\eta'} - P^{\eta\eta'} (q, i\Omega) U^{\eta}]^{-1}.$$
paper we assume that all summations are properly normalized. Summations over momenta \( \sum_{k/q} \) include implicitly a prefactor \( 1/N_{k/q} \) with \( N_{k/q} \) the number of \( k \)-points in the Brillouin zone, and summations of frequencies \( \sum_{i\omega/n\Omega} \) contain implicitly a prefactor \( 1/\beta \) with \( \beta \) the inverse temperature. The SU(2)-broken TRILEX \( \Lambda^2 \) approximation for a single-orbital model reads (cf. Fig. 4 for a diagrammatic representation)

\[
\Sigma_{\beta\alpha}(k, i\omega) = -\sum_{q, i\Omega, \eta} m^\eta \Lambda_{\text{imp}, \alpha}(i\omega + i\Omega, -i\Omega) \tilde{G}_{\beta\eta}(q, i\omega + i\Omega) W_{\text{imp}, \alpha}(i\omega, i\Omega) + \Sigma_{\text{imp}, \beta}(i\omega) \delta_{\alpha\beta} 
\]

\[
P_{\beta\alpha}^\eta(q, i\Omega) = \sum_{k, i\omega, \sigma} \Lambda_{\text{imp}, \alpha}(i\omega + i\Omega, -i\Omega) \tilde{G}_{\beta\eta}(q + k, i\omega + i\Omega) \tilde{G}_{\eta\alpha}(q, i\omega + i\Omega) \Lambda_{\text{imp}, \beta}(i\omega, i\Omega) + P_{\text{imp}, \beta}^\eta(i\Omega) \delta_{\alpha\beta} 
\]

with \( m^\eta = \begin{cases} 2, & \text{if } \eta = xy \\ 1, & \text{else} \end{cases} \)

\[
\sigma' = \begin{cases} \sigma, & \text{if } \eta = xy \\ \text{else}, & \text{else} \end{cases} 
\]

Here, \( \Sigma_{\text{imp}} \) and \( P_{\text{imp}} \) are the self-energy and polarization of the impurity, while \( \alpha \) and \( \beta \) denote sublattice indices (i.e., atoms in the unit cell). Quantities with a tilde in Eq. (24) and (25) are purely non-local, i.e.

\[
\tilde{X}_{\alpha\beta}(k, i\omega) = X_{\alpha\beta}(k, i\omega) - \sum_k X_{\alpha\alpha}(k, i\omega) \delta_{\alpha\beta}. 
\]

In order to be consistent with our choice of the impurity, we have defined locality here on the level of the single atom, instead of the unit cell. Note that here, different from the original TRILEX approximation [51, 52], the separation of local (impurity) and non-local part of the self-energy and polarization is not just a numerical trick to improve on the convergence of frequency summations, but is necessary in order to avoid double counting of local diagrams. This also makes \( \Sigma_{\text{loc}}/P_{\text{loc}} \) trivially equal to \( \Sigma_{\text{imp}}/P_{\text{imp}} \).

In contrast to (E)DMFT, both the fermionic self-energy and the bosonic polarization, acquire a momentum dependence despite the local approximation of the electron-boson coupling vertex of TRILEX. The expressions for the fermionic and bosonic Weiss fields \( G_{\alpha}^\sigma \) and \( U_{\alpha}^\eta \), as well as the impurity action \( S_{\text{imp}, \alpha} \), are identical to (E)DMFT:

\[
G_{\alpha}^\sigma(i\omega) = \left[ G_{\text{loc}, \alpha}(i\omega) \right]^{-1} + \Sigma_{\text{loc}, \alpha}^\sigma(i\omega) \right]^{-1} 
\]

\[
U_{\alpha}^\eta(i\Omega) = W_{\text{loc}, \alpha}^\eta(i\Omega) \left[ \delta_{\eta\eta'} - P_{\text{loc}, \alpha}(i\Omega) W_{\text{loc}, \alpha}^{\eta\eta'}(i\Omega) \right]^{-1} 
\]

where we have defined

\[
D_{\alpha}^{\sigma\sigma'}(\tau) = \left[ U_{\text{ch}}^\sigma(\tau) - U_{\text{ch}}^{\sigma'} \right] + (-)^{\sigma\sigma'} \left[ U_{\text{ch}}^\sigma(\tau) - U_{\text{ch}}^{\sigma'} \right] + (-)^{\sigma'} U_{\text{ch}}^{\sigma\sigma'}(\tau) + (-)^{\sigma} U_{\text{ch}}^{\sigma\sigma'}(\tau) 
\]

\[
J_{\perp, \alpha}(\tau) = 4 \left[ U_{\text{ch}}^{\sigma}(\tau) - U_{\text{xy}}^{\sigma} \right] + 2i \left[ U_{\text{ch}}^{\sigma}(\tau) - U_{\text{ch}}^{\sigma\sigma'}(\tau) \right] 
\]

\[
s_{\pm} = (n^x \pm i n^y)/2, 
\]

using the convention

\[
(-)^{\sigma\sigma'} = \begin{cases} +1 & \text{if } \sigma = \sigma' \\ -1 & \text{if } \sigma \neq \sigma' \end{cases} 
\]

Note the difference between \( U_{\text{xy}}^{\sigma} \), which denotes the two equivalent channel-diagonal elements \( U^{xy} \) and \( U^{yx} \), and \( U^{xy} \) and \( U^{yx} \), which denote channel-off-diagonal contributions. In Eq. (29) we have separated the bare Hubbard interaction \( U \) from the dynamical \( D_{\alpha}^{\sigma\sigma'}(\tau) \).

The TRILEX self-consistency loop now consists of the following steps:

**TR1** Choose a suitable **initial self-energy** \( \Sigma^\sigma(k, i\omega) \) and **polarization** \( P^\eta(q, i\Omega) \).
Use the Dyson equations to calculate the renormalized fermionic and bosonic lattice Green’s functions (Eqs. (19) and (20)).

Calculate the Weiss fields from the local quantities (Eq. (27) and (28)).

Solve the impurity models (for every atom) using the impurity action in Eq. (29). Calculate $\Sigma_{\text{imp}}$, $P_{\text{imp}}$, and $\Lambda_{\text{imp}}$.

Calculate the lattice self-energy and polarization via the Hedin Eqs. (24) and (25).

Inner TRILEX self-consistency loop by repeatedly applying TR2 and TR5 (until convergence).

Go back to TR2.

Note that TR6 is optional. In the highly correlated regime this step however helped stabilizing a paramagnetic solution.

For a more detailed discussion the reader is referred to [51, 52].

In order to solve the paramagnetic KMH model we will also omit a broken sublattice symmetry by solving just one impurity problem, setting the individual impurity quantities equal, i.e.

\[
\Lambda_{\text{imp},\alpha}^{\sigma\eta} = \Lambda_{\text{imp}}^{\sigma\eta},
\]

\[
\Sigma_{\text{imp},\alpha}^\sigma = \Sigma_{\text{imp}}^\sigma,
\]

\[
P_{\text{imp},\alpha}^{\eta\eta'} = P_{\text{imp}}^{\eta\eta'}.
\]

Enforcing paramagnetism then means

\[
\sigma_{\eta\uparrow}\Lambda_{\text{imp}}^{\eta\eta} = \sigma_{\sigma\uparrow}^{\eta\eta} \Lambda_{\text{imp}}^{\eta\eta},
\]

\[
\Sigma_{\text{imp}}^\sigma = \Sigma_{\text{imp}}^\sigma = \Sigma_{\text{imp}}^\sigma.
\]

\[
P_{\text{imp}}^{\eta\eta'} = P_{\text{imp}}^{\eta\eta'} \delta_{\eta\eta'}.
\]

For more details on why Eqs. (36) and (38) hold in the paramagnetic case see Appendix B and C. The self-energy and polarization in our approximation therefore read

\[
\Sigma^\sigma = \left(\frac{\Sigma_{AA}}{\Sigma_{BA}} + \Sigma_{\text{imp}} \delta_{AB} + \Sigma_{\text{imp}} \right)
\]

\[
P_{\eta\eta'}^{\eta\eta'} = \left(\frac{\tilde{P}_{\eta\eta'}^{\eta\eta'}}{\tilde{P}_{\text{imp}}^{\eta\eta'}} + \tilde{P}_{\text{imp}}^{\eta\eta'} \delta_{\eta\eta'} \right).
\]

In Appendix C we also show that $P_{\eta\eta'}^{\text{ch}} (P_{\eta\eta'}^{\text{oc}})$ is the only non-zero off-diagonal contribution to the KMH model, when calculations are performed at half-filling, where the system is particle-hole symmetric [70].

Calculations have been performed using a 30 × 30 k-point grid. The TRILEX ($\Lambda^2$) code has been implemented using the TRIQS library [61], and uses an implementation of Rubtsov’s interaction expansion continuous-time quantum Monte-Carlo algorithm [71] as an impurity solver.

In [69] also paramagnetic calculations are performed on a model with non-local SOC using a dual-boson based approach. Quite generally, the TRILEX $\Lambda^2$ method presented here is related to other beyond-DMFT schemes such as the dual-boson [72, 73] or the D-TRILEX approach [66, 68, 74]. In all these approaches a double insertion of the vertex is performed. In the dual-boson schemes, since local and non-local contributions are separated, there is no problem with double counting of diagrams. The double insertion of the vertex just leads to more diagrams that are taken into account. Let us stress here that dual-boson methods have been designed to target non-local interactions, while in our work we are restricting ourselves to local interactions only.

A. Determination of the $\alpha$-parameter

The choice of the Fierz parameter $\alpha$ in Eq. (17) is influencing the results both at the one- and two-particle level [51, 75]. This so called Fierz-ambiguity can be systematically reduced when the electron-boson vertex is extracted from clusters of increasing size instead of single-site impurities [64], however at significantly increased computational costs.

For the two dimensional square lattice close to the antiferromagnetic (AF) phase transition (at $T = 0$) spin-fluctuations are dominant [76]. As we are enforcing a paramagnetic solution also in an AF regime, all calculations in the main text will refer to $\alpha = 1/3$ [75], unless stated otherwise. In order to demonstrate that our conclusions do not depend on the choice of $\alpha$, in Appendix E we show results for a wide range of Fierz parameters.

V. RESULTS

To show that local interactions increase the effective spin-orbit coupling in the KMH model, we first take a look at the structure of $\Sigma^{SO}$ and see if it matches the structure of the spin-orbit term of the Hamiltonian. Therefore, we perform a Fourier transformation to real space and compare $\text{Im} \Sigma^{SO}(r, i\omega_0)$ to the SOC term in Eq. (1). From Fig. 5 we see that just like in Eq. (1) pre-dominantly nearest-neighbor terms contribute to $\Sigma^{SO}$, and that right turns pick up a different sign than left turns. The real part turns out to be negligible. This relationship also holds for higher Matsubara frequencies,
From Eq. (4) we see that $k$ to (41c), in next-nearest neighbor hopping term in real-space (Eq. SO $k$ self-energy, we take its representation in (see also Fig. 2). For the analytic continuation of the with $a$ the system would order above a critical $U$ the system would order above a critical $\lambda_{eff} = 7U$ [34] was found. The authors $U$ and $\alpha = 0.33$. $\lambda_{eff}^{1B}$ increases gently.

At this point we want to emphasize again that we are performing paramagnetic calculations by symmetrizing the impurity quantities. Without symmetrization the system would order above a critical $U^{c}(AF)$ between $4t - 5t$, depending on the underlying method. In lattice quantum Monte Carlo a value around $U^{c} \gtrsim 5t$ [25], and for cluster-DMFT $U^{c} \sim 5t$ [34] was found. The authors find $U^{c} \gtrsim 4t$ for single-site DMFT. This means that half of the plot in Fig. 7 belongs to a regime that has a high tendency towards ordering. We had to use a quite low mixing factor (between 0.3 and 0.15) of the Weiss fields (TR3) but also of the lattice self-energy and polarization in the inner TRILEx self-consistency loop (TR6) in order to stabilize paramagnetic solutions. Calculations with $U = 7.5t$ have become already quite unstable (denoted by the dashed line in Fig. 7 and 9) and we have not been able to stabilize solutions for larger values of $U$. Note however that enforcing the paramagnetic solu-

\[
\text{Im} \Sigma^{SO}(r - r' = a_{ij}, i\omega) \propto \nu_{ij} \lambda_{SO}^{1B} \forall i\omega, \quad (41a)
\]
\[
\left| \text{Im} \Sigma^{SO}(r - r' \neq a_{ij}, i\omega) \right| < 0.2 \forall i\omega, \quad (41b)
\]
\[
\left| \text{Im} \Sigma^{SO}(r - r' = a_{ij}, i\omega) \right| < 2 \cdot 10^{-3} \forall i\omega, \quad (41c)
\]

with $a_{ij}$ being a primitive unit vector. This shows that $\Sigma^{SO}$ has the same structure as the SOC term in Eq. (1) (see also Fig. 2). For the analytic continuation of the self-energy, we take its representation in $k$-space. According to the correspondence of $\Sigma^{SO}$ to the complex next-nearest neighbor hopping term in real-space (Eq. (41a) to (41c)), in $k$-space we find

\[
\left| \text{Im} \Sigma^{SO}(k, i\omega) \right| < |\text{Re} \Sigma^{SO}(k, i\omega)| \forall i\omega. \quad (42b)
\]

From Eq. (4) we see that $\gamma_{k = K/3} = \mp 3\sqrt{3}\lambda_{SO}^{1B}$ and, hence, together with Eq. (41a) we find

\[
\nu_{ij} \text{Im} \Sigma^{SO}(r - r' = a_{ij}, i\omega) \approx \frac{\text{Re} \Sigma^{SO}(k = K', i\omega)}{3\sqrt{3}}. \quad (43)
\]

Note that this relation becomes exact if the terms in Eq. (41b) and (41c) completely vanish. We can thus analytically continue the right-hand side of Eq. (43) to get the frequency-dependent SOC enhancement [77]. We perform this analytic continuation using the maximum entropy method implemented in the TRIQS library [78, 79]. The result is shown in the lower panel of Fig. 6 for several different values of $U$, and we find that at least for small frequencies the SOC-enhancement is a constant, making our definition of $\lambda_{eff}$ in Eq. (11) reasonable.

In Fig. 7 we show how $\lambda_{eff}^{1B}$ develops with increasing on-site interaction $U/t$. In order to quantify the correlation strength accompanied by the different values of $U/t$, the renormalization factor

\[
Z = \lim_{\omega \to 0} \left[ 1 - \frac{\partial \text{Im} \Sigma^{imp}(i\omega)}{\partial \omega} \right]^{-1} \quad (44)
\]

is also shown in Fig. 7. In order to approximate the derivative in Eq. (44) we use the slope between the two Matsubara frequencies $\omega_0$ and $\omega_1$. This is not a real quasi-particle weight in the sense of a Landau-liquid theorem (there are no states at the Fermi-level in the non-interacting model already, see Fig. 3), however, this still quantifies the level of correlation in the system and is also bound between 1 and 0. From Fig. 7 we see how, while correlations are getting stronger ($Z$ is dropping), $\lambda_{eff}^{1B}$ increases gently.

At this point we want to emphasize again that we are performing paramagnetic calculations by symmetrizing the impurity quantities. Without symmetrization the system would order above a critical $U^{c}(AF)$ between $4t - 5t$, depending on the underlying method. In lattice quantum Monte Carlo a value around $U^{c} \gtrsim 5t$ [25], and for cluster-DMFT $U^{c} \sim 5t$ [34] was found. The authors find $U^{c} \gtrsim 4t$ for single-site DMFT. This means that half of the plot in Fig. 7 belongs to a regime that has a high tendency towards ordering. We had to use a quite low mixing factor (between 0.3 and 0.15) of the Weiss fields (TR3) but also of the lattice self-energy and polarization in the inner TRILEx self-consistency loop (TR6) in order to stabilize paramagnetic solutions. Calculations with $U = 7.5t$ have become already quite unstable (denoted by the dashed line in Fig. 7 and 9) and we have not been able to stabilize solutions for larger values of $U$. Note however that enforcing the paramagnetic solu-
FIG. 7. Evolution of the $\lambda^{1B}_{\text{eff}}/t$ and the renormalization factor $Z$ with increasing interaction $U/t$. The dashed line denotes a regime where it becomes increasingly hard to stabilize a paramagnetic solution.

The definition is important in order to make the one-band model comparable to the two-band model.

For the two-band model of bismuthene we also look at the real-frequency dependence of the SOC-enhancement, again by applying the maximum entropy method. The top panel of Fig. 6 shows that also here the enhancement is roughly constant in frequency. Unlike the one-band model this holds even at large frequencies. Hence the definition of $\lambda^{2B}_{\text{eff}}$ in Eq. (14) is also plausible. Fig. 8 shows that for bismuthene $Z$ decreases quite linearly with $U/t$, while $\lambda^{2B}_{\text{eff}}$ starts diverging.

In order to compare the one-band and two-band model, Fig. 9 shows for both models the relative increment of SOC $\lambda_{\text{eff}}/\lambda_{SO}$ as a function of the renormalization factor $Z$. Here we have further added results for additional $\alpha$-parameters in the interval $[1/3,1/2]$, showing that results remain qualitatively consistent over the full range of $\alpha$-values (c.f. also Appendix E 1). We find that the enhancement of non-local SOC in the one-band model is negligible compared to the enhancement of local SOC in the two-band model.

The reason for this different response of the effective SOC to interactions can be traced back at least qualitatively to the specific forms of the SOC Hamiltonians as compared to the interaction Hamiltonians. For the one-band model with a non-local SOC Hamiltonian, the effective SOC is defined from a non-local self-energy, while the microscopic Hubbard interaction is purely local. For the two-band model, however, where the relevant self-energy component for the effective SOC is a local but orbital off-diagonal one, the local microscopic Kanamori interaction includes corresponding inter-orbital interaction terms. Thus, the interaction Hamiltonian is directly effecting the aforementioned self-energy terms. We believe this discrepancy between the locality of the SOC and the microscopic interaction to be the reason for the stronger response in the two-band case as opposed to the one-band case. In order to corroborate this assumption, we can treat the effective enhancement of SOC in Hartree-Fock (HF) theory. As we detail in Appendix D, the HF decoupling in the two-band case leads to a contribution that is exactly of the form of the SOC Hamiltonian. For the one-band case, such a contribution is absent by construction.

We can also quantify the HF estimate for the SOC enhancement. First, we evaluate the SOC formula for the enhancement (Eq. (D1)), using the converged DMFT results for the spin-orbital polarization, which is shown as green line in the lower panel of Fig. 8. Second, we also solved the HF equations self-consistently, the result of which is shown as the purple line in the above mentioned figure. In general, both HF estimates overestimate the DMFT results, where the self-consistent solution shows particularly strong enhancement. The reason is that in HF, already a rather small interaction $U$ and $J$ is sufficient to give an almost saturated polarization, and in
turn strong enhancement. This is consistent with the considerations in Ref. [11], where from an almost fully spin-orbital polarized system a rather small interaction $U - J = 0.5$ eV was deduced. As a result, these HF estimates show that it is important to take the dynamical correlation effects in the system properly into account.

VI. CONCLUSIONS

In this work we investigated the effective SOC strength in a one-band and a two-band honeycomb lattice model. The main difference between the models is the structure of the SOC coupling term, which in the one-band case is non-local as opposed to the purely local term in the two-band case. In order to quantify this effect, we have defined an effective SOC $\lambda_{1B/2B}^{\text{eff}}$ (Eqs. (11) and (14)). Both increase with increasing interaction strength. However, the effective enhancement in the one-band case is rather minute. One main reason for this difference is that in the two-band model both the interactions and the SOC are local terms. We find that a Hartree-Fock treatment of the interactions already leads to a sizable enhancement of the SOC. This effect is completely absent in the one-band model. However, we have furthermore shown that the Hartree-Fock treatment alone drastically overestimates this enhancement.

For the two-band case, within the treatment of DMFT, we see strong enhancements up to factors of 3 when correlations become sizable. Thus, for a similar correlation strength as measured by the renormalization factor $Z$, the enhancement in the two-band model is approximately one order of magnitude larger than in the one-band model. This suggests that effective SOC is a relevant concept in multi-orbital systems, but might be of rather limited importance in single-band systems with non-local SOC, at least for the correlation regime that we studied here. This possibly changes in the vicinity of phase transitions, or when also non-local interactions are included into the KMH, as they may considerably enhance the non-local effective SOC.

On the technical level, we have formulated the necessary equations for TRILEX in the case when SU(2) symmetry is broken. Since the original TRILEX formulations break hermiticity of certain quantities, we adopted the so-called TRILEX $\Lambda^2$ method. This broken symmetry is not only relevant for the study that we presented here, but of course also for models with long-range magnetic ordering. The formalism, without the enforced paramagnetic properties as developed here, is directly applicable to the magnetic case.

ACKNOWLEDGMENTS

We thank T. Ayral and O. Parcollet for insightful discussions. This work has been supported by the Austrian Science Fund (FWF), grant number Y746. A part of the calculations have been performed on the Vienna Scientific Cluster.

Appendix A: Derivation of the single-orbital SU(2)-broken TRILEX equations

The most general formulation of the Hedin equations for the fermionic self-energy and the bosonic polarization reads [52]

\[
\Sigma_{ji}^{\sigma} = \sum_{\eta} \lambda_{aib}^{\sigma\eta} G_{ai}^\sigma G_{bj}^{\eta'} \Lambda_{ba}^{\eta\eta'} \quad \text{(A1)}
\]

where $a, b, c, d, e, f, i, j$ are space-time indices, $\sigma, \sigma', \tilde{\sigma}, \tilde{\sigma}'$ denote spin indices and $\eta, \eta'$ denote Pauli indices $\{0, x, y, z\}$ and Einstein’s summation convention is adopted. $\lambda$ is the bare fermion-boson coupling vertex and $\Lambda$ its fully renormalized version.

For the derivation of the equations for the specific purpose of this paper, we assume a spin-diagonal form of the fermionic Green’s functions, i.e. $G_{ij}^{\sigma\sigma'} = G_{ij}^{\sigma\delta_{\sigma\sigma'}}$. The polarization hence becomes

\[
P_{ji}^{\eta\eta'} = \lambda_{aib}^{\sigma\eta} G_{ai}^\sigma G_{bj}^{\eta'} \Lambda_{ba}^{\eta\eta'} \quad \text{(A2)}
\]

For the vertex $\Lambda_{ba}^{\eta\eta'}$ spin and channel indices are connected in the sense that $\Lambda_{ba}^{\sigma\eta} = \Lambda_{ba}^{\eta\sigma}|_{\sigma^*\sigma}$. Given that, we find

\[
P_{ji}^{\eta\eta'} = \lambda_{aib}^{\sigma\eta} G_{ai}^\sigma G_{bj}^{\eta'} \Lambda_{ba}^{\eta\eta'} \quad \sigma' = \begin{cases} \{0, x, y\}, & \text{if } \eta, \eta' \in [0, x, y] \\ \{0, x, y\}, & \text{if } \eta, \eta' \in [0, x, y] \end{cases}
\]

(A3)

which is Eq. (21) of the main text. This gives, besides, the diagonal terms $P_{ji}^{\eta\eta'}$ also 4 non-zero off-diagonal terms: Using the same assumptions as above for the self-energy we find

\[
\Sigma_{ji}^{\tilde{\sigma}\tilde{\sigma'}} = -\lambda_{aib}^{\tilde{\sigma}\eta} G_{ai}^\sigma G_{bj}^{\eta'} \Lambda_{ba}^{\eta\eta'} \quad \text{(A4)}
\]
and hence, by explicitly writing the vertex
\[
\Sigma^{\bar{\sigma}\sigma}_{ji} = -\lambda^{\sigma\eta}_{aib} G_{d\sigma}^\sigma W_{bc}^{\eta\eta'} \Lambda_{jdc}^{\sigma'\eta'} |\sigma\rangle |\sigma'\rangle. 
\] (A5)

Given a density-density interaction as presented in Eq. (15), the bare part of \( W^{\eta\eta'} \) will be fully channel-diagonal, and channel-offdiagonal components get generated only through the polarization (see Eq. (20)). As \( P \) is block-diagonal (\([ch,z] \) and \([x,y] \)) the restrictions of Eq. (A3) pass on to \( W^{\eta\eta'} \). Hence we only get non-zero contributions to the self-energy if \( \sigma = \bar{\sigma} \), leaving \( \Sigma_{ij} \) spin-diagonal. The equation simplifies to
\[
\Sigma^{\bar{\sigma}\sigma}_{ji} = -\lambda^{\sigma\eta}_{aib} G_{d\sigma}^\sigma W_{bc}^{\eta\eta'} \Lambda_{jdc}^{\sigma'\eta'} |\bar{\sigma}\rangle |\eta\rangle, 
\] (A6)

which is Eq. (22) of the main text.

The graphical representations of Eqs. (A3) and (A6) can be found in Fig. 10.

1. **Local vertex approximation**

a. **TRILEX**

Now we will use the fact that we are only considering Hubbard-type interactions. Therefore the bare vertex can be rewritten as \( \lambda^{\sigma\eta}_{aib} = \sigma_{\bar{\sigma}\sigma}\delta_{ai}\delta_{ib} \) and Eq. (A6) becomes
\[
\Sigma^{\bar{\sigma}\sigma}_{ji} = -\sigma_{\bar{\sigma}\sigma}\lambda_{imp,\bar{\sigma}} \Lambda_{jdc}^{\sigma'\eta'} \] (A7)

In the following \( \alpha, \beta, \gamma, \delta \) denote inner unit cell indices (e.g. \( A,B \)) and \( R,R_1,R_2 \) are lattice vectors connecting unit cells. By omitting the spin/channel indices for clarity Eq. (A6) becomes:

\[
\Sigma_{\beta\alpha}(R,\tau) = -\int_{\tau_1,\tau_2} \sum_{R_1,R_2,\gamma,\delta} G_{\gamma\alpha}(R_1,\tau_1) W_{\alpha\delta}(-R_2,\tau_2) \Lambda_{\beta,\gamma,\delta}(R-R_1,R_2-R_1,\tau-\tau_1,\tau_2-\tau_1). \] (A8)

Now, assuming the locality of the electron-boson coupling vertex
\[
\Lambda_{\beta,\gamma,\delta}(R-R_1,R_2-R_1,\tau-\tau_1,\tau_2-\tau_1) \approx \Lambda_{\beta}(\tau-\tau_1,\tau_2-\tau_1) \delta_{\beta,\gamma,\delta} \delta_{RR_1}\delta_{RR_2}, \] (A9)

and approximating the local vertex by the impurity vertex (TRILEX approximation)
\[
\Lambda_{\beta} = \Lambda_{imp,\beta}, \] (A10)

we obtain
\[
\Sigma_{\beta\alpha}(R,\tau) = -\int_{\tau_1,\tau_2} G_{\gamma\alpha}(R,\tau_1) W_{\alpha,\beta}(-R_2,\tau_2) \Lambda_{imp,\beta}(\tau-\tau_1,\tau_2-\tau_1). \] (A11)

Taking the following definition of the Fourier transform
\[
X_{\alpha\beta}(k,i\omega) = \int_{\tau} e^{ikR\tau} X_{\alpha\beta}(R,\tau), \] (A12)

we find by also adding back spin/channel indices
\[
\Sigma_{\beta\alpha}(k,i\omega) = -\sum_{q,i\Omega} \sigma^{\gamma\eta}_{\bar{\sigma}\sigma} G_{\beta\alpha}^{\gamma\eta} (k+q,i\omega+i\Omega) W_{\alpha,\beta}^{\eta\eta'} (k,i\omega) \Lambda_{\gamma,\eta,\delta}^{\eta',\gamma'} (i\omega,i\Omega) \] (A13)

and equivalently
\[
P_{\beta\alpha}^{\eta\eta'} (q,i\Omega) = \sum_{k,i\omega,\sigma} \sigma^{\gamma\eta}_{\bar{\sigma}\sigma} G_{\alpha\beta}^{\gamma\eta} (k+q,i\omega+i\Omega) G_{\alpha\beta}^{\eta\eta'} (k,i\omega) \Lambda_{\gamma,\eta,\delta}^{\eta',\gamma'} (i\omega,i\Omega). \] (A14)

This formulation does not enforce the required Hermitian symmetry for channel-off-diagonal elements namely \( P_{\alpha\beta}^{\eta\eta'} (q,i\Omega) = [P_{\beta\alpha}^{\eta\eta'} ((q,-i\Omega))]^* \). In order to stay consistent it is necessary to restrict the bosonic Green’s functions to
\[
X_{\eta\eta'}^{\dagger} = X_{\eta\eta'} \delta_{\eta\eta'}. \] (A15)
b. TRILEX $\Lambda^2$

In order to enforce hermiticity of the polarization without requiring channel-diagonality we adopt the TRILEX $\Lambda^2$ approach, where the non-local part of the self-energy $\tilde{\Sigma}$ and polarization $\tilde{P}$ is calculated from a diagram with two renormalized impurity-vertices appearing on both sides. The non-local part of Eq. (A11) hence becomes

$$\tilde{\Sigma}_{\beta\alpha}(\mathbf{R}, \tau) = -\int_{\tau_1, \tau_2, \tau_3, \tau_4} \Lambda_{\text{imp}, \alpha}(\tau_1, \tau_2) \tilde{G}_{\beta\alpha}(\mathbf{R}, \tau - \tau) \tilde{W}_{\alpha, \beta}(\mathbf{R}, \tau_2 - \tau_4) \Lambda_{\text{imp}, \beta}(\tau - \tau_3, \tau_4 - \tau_3).$$

(A16)

and after Fourier transform we obtain

$$\tilde{\Sigma}_{\beta\alpha}(\mathbf{k}, i\omega) = -\sum_{\mathbf{q} \in RBZ, i\Omega} \Lambda_{\text{imp}, \alpha}(i\omega + i\Omega, -i\Omega) \tilde{G}_{\beta\alpha}(\mathbf{q} + \mathbf{k}, i\omega + i\Omega) \tilde{W}_{\alpha, \beta}(\mathbf{q}, i\Omega) \Lambda_{\text{imp}, \beta}(i\omega, i\Omega),$$

(A17)

and similarly

$$\tilde{P}_{\beta\alpha}(\mathbf{q}, i\Omega) = \sum_{\mathbf{q} \in RBZ, i\Omega} \Lambda_{\text{imp}, \alpha}(i\omega + i\Omega, -i\Omega) \tilde{G}_{\beta\alpha}(\mathbf{q} + \mathbf{k}, i\omega + i\Omega) \tilde{G}_{\alpha, \beta}(\mathbf{k}, i\omega) \Lambda_{\text{imp}, \beta}(i\omega, i\Omega).$$

(A18)

Recovering spin and channel indices and replacing the local quantities $\Sigma_{\text{loc}}/P_{\text{loc}}$ by the impurity ones $\Sigma_{\text{imp}}/P_{\text{imp}}$, leads Eq. (24) and (25) in the main text.

2. Calculation of the Vertex

The full electron-boson coupling vertex can be written as [52]

$$\Lambda_{\alpha\beta\gamma}^{\sigma\sigma'} = \left[G^{-1}\right]_{\alpha\gamma}^{\sigma'} \left[G^{-1}\right]_{\beta\gamma}^{\sigma} \left[\delta_{\sigma\sigma'} - \chi_{\alpha\beta\gamma}^{\eta\eta'} U_{\alpha i} \right]^{-1} \chi_{\alpha\beta\gamma}^{\eta\eta'},$$

(A19)

with the fermionic three-point correlation function

$$\chi_{\alpha\beta\gamma}^{\eta\eta'} = \left\langle \hat{c}_{\alpha} \hat{c}_{\beta} \left( n_{\gamma}^{\eta} - \left\langle n_{\gamma}^{\eta} \right\rangle \right) \right\rangle,$$

(A20)

the susceptibility $\chi_{\alpha\beta\gamma}^{\eta\eta'}$ (see Eq. (C1)) and the decoupled interaction $U_{\alpha i}^{\eta}$ (see Eq. (15)). Here again $a - i$ are space-time indices, $\hat{c}_{\alpha}^{\dagger}$ ($\hat{c}_{\alpha}$) are creation (annihilation) operators and $n_{\gamma}^{\eta}$ is given by Eq. (16). Applying the same considerations as above and replacing the interaction by the according Weiss field (Eq. (28)), the impurity vertex is given by

$$\Lambda_{\text{imp}, \alpha}^{\sigma\eta}(i\omega, i\Omega) = \left[G_{\text{imp}, \alpha}^{\sigma'}(i\omega + i\Omega)\right]^{-1} \left[G_{\text{imp}, \alpha}^{\sigma}(i\omega)\right]^{-1} \left[\delta_{\sigma\sigma'} - \chi_{\text{imp}, \alpha}^{\eta\eta'}(i\Omega) U_{\alpha i}^{\kappa \kappa'}(i\Omega)\right]^{-1} \chi_{\text{imp}, \alpha}^{\eta\eta'}(i\omega, i\Omega).$$

(A21)

Appendix B: Implementation details for the paramagnetic KMH

Calculating the impurity vertex for a single-site paramagnetic calculation (see Eq. (33) and (36)) reduces Eq. (A21) to

$$\Lambda_{\text{imp}}^{\eta\eta}(i\omega, i\Omega) = \frac{\chi_{\text{imp}}^{\eta\eta}(i\omega, i\Omega)}{G_{\text{imp}}^{\eta\eta}(i\omega + i\Omega) G_{\text{imp}}^{\eta\eta}(i\omega) (1 - \chi_{\text{imp}}^{\eta\eta}(i\Omega) U_{i}^{\eta\eta}(i\Omega))},$$

(B1)

Note that Eq. (B3) directly follows from rewriting the local version of (A20) in terms of creation and annihilation-operators only and by requiring, similar to Eq. (C5b), all terms with flipped spins to be equal. As there is no other spin-dependency in Eq. (B1) but $\chi_{\text{imp}}$, Eq. (36) directly follows.
that there are time-ordering operator $T$ with space indices $\alpha$ and $\eta$, namely Hartree-Fock and second-order perturbation theory.

In the following we apply two analytic approaches, namely Hartree-Fock and second-order perturbation theory.
FIG. 13. Frequency dependence of SOC-enhancement of the 1-band model (lower figure) for $U = 7.0t$ (like Fig. 6) for different $\alpha$.

1. Hartree-Fock

For a two-band model the Hartree-Fock decoupling gives a contribution to the effective SOC as [11]

$$\lambda_{\text{eff}}^{2B,\text{HF}} = \lambda_{\text{SO}}^{2B} + \frac{1}{2}(U - J)\left(\langle n_{\uparrow}^\dagger \rangle - \langle n_{\downarrow}^\dagger \rangle\right), \quad (D1)$$

where $n_{\sigma}^\dagger$ are density operators in the eigenbasis (Eq. (12)) of the spin-orbit coupling Hamiltonian $H_{\text{SO}}$. This contribution is derived from the full interaction Hamiltonian in Hartree-Fock approximation given by

$$H_{\text{int, HF}}^{2B} = \sum_{\sigma} [(U - 2J)d^\sigma + (2U - 3J)d^\sigma^\dagger] (n_{\sigma}^\dagger + n_{\sigma}), \quad (D2)$$

where we have defined $d^\sigma = \langle n_{\sigma}^\dagger \rangle = \langle n_{\sigma}^\dagger \rangle$ in order to enforce paramagnetism.

The enhancement given by Eq. (D1) with densities extracted from the DMFT solution, must however be equal to the high-frequency ($i\omega \to \infty$) part of the related self-energy components. Hence, comparing Eq. (D1) to (14), the relation

$$\frac{(U - J)}{2} \left(\langle n_{\uparrow}^\dagger \rangle - \langle n_{\downarrow}^\dagger \rangle\right) = \text{Re} \left(\Sigma_{\uparrow}(i\omega_\infty) - \Sigma_{\downarrow}(i\omega_\infty)\right)$$

must hold, which we can confirm for all calculations presented here.

In Fig. 8 we compare the self-consistent Hartree-Fock solution to the high- and low-frequency contribution of DMFT and find that Hartree-Fock highly overestimates the enhancement. This is consistent with the fact that there is already a big difference between the high- and low-frequency contribution of DMFT. This difference is illustrated in Fig. 14. Note that we have plotted the frequency-dependence of the enhancement on the imaginary axis, as the analytic continuation becomes increasingly complicated with higher frequencies. Nevertheless, the values at the two points $i\omega \to 0$ and $i\omega \to \infty$ are the same on real and imaginary axis.

It is easily seen by looking at Eq. (1), that the Hartree-Fock approximation cannot give a contribution to the spin-orbit coupling of the one-band model, as the interaction and the SOC term do not involve the same bonds.

FIG. 14. Frequency dependence of the SOC-enhancement for the two-band model for $U = 6.7t$ and $J = 1.3t$ on the imaginary axis. The high-frequency contribution is calculated from Eq. (D3).

FIG. 15. $2^{nd}$ order expansion for single-site model with onsite-interaction only. $G_0^\sigma$ is the non-interacting Green’s function and $U$ the bare interaction.

All we get is the term $U \langle n_{i\sigma} \rangle$. For a paramagnetic calculation this gives the same constant for every spin (and site) and hence gives only a contribution to the chemical potential. Therefore the Hartree-Fock estimate of the effective SOC is given by the bare SOC:

$$\lambda_{\text{eff}}^{1B,\text{HF}} = \lambda_{\text{SO}}^{1B} \quad (D4)$$

This is in agreement with the high frequency behavior of our TRILEX solution.

2. Second order perturbation theory

In order to get a non-vanishing contribution to the SOC also in the one band model, we need to include further contributions to the self-energy. Therefore we have a look at the second order contribution to the self-energy. For a single orbital model with only onsite-interaction like the KMH-model this is given by

$$\Sigma_{\sigma, 2^{nd}}(\mathbf{r}, \tau) = -U^2 G_0^\sigma(\mathbf{r}, \tau) G_0^\sigma(\mathbf{r}, \tau) G_0^\sigma(-\mathbf{r}, -\tau. \quad (D5)$$

where $G_0^\sigma$ is the non-interacting propagator. The according diagrams are shown in Fig. 15. Calculating the contribution to the SOC again as described in sec. III A, for $\lambda_{\text{SO}}^{1B} = 0.14t$ we find

$$\lambda_{\text{eff}}^{1B, 2^{nd}} = 0.14t + 6.8 \cdot 10^{-5} U^2 t, \quad (D6)$$

which leads an enhancement of the SOC-coupling that is roughly one order of magnitude smaller than we observe by performing TRILEX calculations. However also
this study qualitatively shows that the effective SOC increases when correlations are included.

Appendix E: Some more details on the self-energy

1. Fierz-ambiguity of $\lambda_{\text{eff}}^{1B}$

As mentioned in subsection IV A, the choice of the decoupling parameter $\alpha$ can influence the results, especially when single-site impurity problems are solved. In Fig. 12 we show the development of $\lambda_{\text{eff}}^{1B}$ and $Z$ with increasing local interaction $U$ for different decoupling parameters $\alpha$. We find that while the renormalization factor is affected only weakly by the choice of $\alpha$, $\lambda_{\text{eff}}^{1B}$ decreases in a monotonic fashion when we increase the Fierz parameter from a pure spin decoupling ($\alpha = 1/3$) to a pure charge decoupling ($\alpha = 2/3$). The calculation for $\alpha = 2/3$ turned out to be unstable and, therefore, is not contained in our results, but the tendency becomes clear already for values of $\alpha$ below $2/3$. The qualitative results of the main text remain unchanged, as all calculations for $\alpha > 1/3$ lead to lower effective SOC (see also Fig. 9). We can also see from Fig. 12 how the dependence of the results on the Fierz-parameter is smaller for TRILEX $\Lambda$ than for TRILEX. Mind, however, that we are also ignoring channel-off-diagonal components in the TRILEX approach (see Eq. (A15)) that are small but not vanishing (see Fig. 11).

We note in passing that it has been argued that the Fierz-ambiguity can be resolved within the dual TRILEX approach [66].

2. Frequency-dependence of SOC-enhancement

For the sake of completeness we also look at the frequency dependence of the SOC-enhancement for different Fierz-parameter, i.e., we did analytic continuations of $\text{Re} \Sigma^\text{SO}$ as we did in the main text, but for different $\alpha$ parameters, which is shown in Fig. 13. We see clearly that all calculations for different $\alpha$ follow a very similar trend, namely a constant value for small frequencies followed by a small upturn.

3. Effect of SOC on the normal non-SOC part of the self-energy

An interesting question is how the inclusion of SOC changes the normal, i.e. time-reversal symmetry preserving part of the self-energy. In the two-band model, this symmetry preserving part is given by $\frac{1}{2} (\Sigma^{\uparrow}_d + \Sigma^{\uparrow}_d)$. For the one-band model, both $\Sigma^d$ and $\Sigma_{AB}$ are the normal components. In order to make a comparison between the two models we will, however, solely focus on the impurity self energies, which for the one-band case is $\Sigma_{\text{imp}} = \sum_k \Sigma^d$.

Fig. 16 shows the effect of introducing a SOC-term in both models. We can see that in either case the correlations are reduced when $\lambda_{\text{SO}}$ is turned on. However, for the two-band model for bismuthene this effect is much stronger than for the KMH model. This goes hand in hand with a much larger impact of correlations on the effective SOC in the two-band model, as discussed in the main text. Obviously, here the inversion of the arguments is true, namely that not only correlations have a strong impact on the effective SOC, but also that SOC has a strong impact on correlations related to normal components of the self-energy. We notice in passing that the choice of the decoupling parameter $\alpha$ has a rather small quantitative impact on this qualitative result.

a. $k$-dependence of $\Sigma^d$

Last but not least we want to look at the effect of $\lambda_{\text{SO}}^{1B}$ on the $k$-dependence of $\Sigma^d$. From Fig. 17 we find that SOC reduces the $k$-dependence of $\text{Im} \Sigma^d$ and hence again the strength of the correlation. In order to show the development of the $k$-dependence with increasing interactions we introduce $\text{Im} (\Sigma^d_\Gamma(i\omega_n) - \Sigma^d_K(i\omega_n))$ as a measure for the non-locality of the self-energy. We consider this reasonable as it involves the two most different values in the Brillouin zone. Fig. 18 shows this measure for differ-
FIG. 17. Im $\Sigma^d(k, i\omega_0)$ at $U = 4.8t$ and $\alpha = 0.33$ for $\lambda^{1B}_{SO} = 0$ (upper figure) and $\lambda^{1B}_{SO} = 0.14t$ (lower figure).

FIG. 18. Difference of the two most distant points in Im $\Sigma^d(k, i\omega_0)$ $\Gamma$ and $K$ as a measure of the $k$-dependence of $\Sigma^d$ for $\lambda^{1B}_{SO}$ and different parameters of $U$ and $\alpha$. We again compare TRILEX $\Lambda^2$ to TRILEX as in Fig. 12.

ent interaction values $U$ and decouplings $\alpha$. Not surprisingly, the $k$-dependence increases with increasing $U$, as correlations become stronger. Contrary to Fig. 16, but similar to Fig. 12, we find that the relative effect of $\alpha$ on the $k$-dependence is clearly visible in this measure. This again shows that for the KMH model non-local quantities are more affected by the choice for the decoupling parameter than local quantities. However, we again find that the treatment in TRILEX $\Lambda^2$ decreases the Fierz-ambiguity as compared to TRILEX.

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