Semirealistic tight-binding model for spin-orbit torques

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We compute the spin-orbit torque in a transition-metal heterostructure using Slater-Koster parametrization in the two-center tight-binding approximation and accounting for d orbitals only. In this method, the spin-orbit coupling is modeled within Russell-Saunders scheme, which enables us to treat interfacial and bulk spin-orbit transport on equal footing. The two components of the spin-orbit torque, dissipative (dampinglike) and reactive (fieldlike), are computed within Kubo linear response theory. By systematically studying their thickness and angular dependence, we were able to accurately characterize these components beyond the traditional inverse spin galvanic and spin Hall effects. We find that the conventional fieldlike torque is purely interfacial. In contrast, we unambiguously demonstrate that the conventional dampinglike torque possesses both an interfacial and a bulk contribution. In addition, both fieldlike and dampinglike torques display substantial angular dependence with strikingly different thickness behaviors. While the planar contribution of the fieldlike torque decreases smoothly with the nonmagnetic metal thickness, the planar contribution of the dampinglike torque increases dramatically with the nonmagnetic metal thickness. Finally, we investigate the self-torque exerted on the ferromagnet when the spin-orbit coupling of the nonmagnetic metal is turned off. Our results suggest that the spin accumulation that builds up inside the ferromagnet can be large enough to induce magnetic excitations.

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I. INTRODUCTION

Current-driven spin-orbit torques have become a prodigal area of research in the past ten years [1,2]. This magnetic torque enables the electrical control of ferromagnets using spin densities generated through angular momentum transfer between the orbital and spin degrees of freedom [3,4]. Understanding the physical origin of the torques, their symmetries and materials dependence has been the subject of intense collaborations between experimentalists and theorists. From the theory standpoint, several mechanisms have been identified, among them spin Hall effect [5], inverse spin galvanic effect [6–8] (also called Rashba-Edelstein effect), spin swapping [9–11], interfacial spin precession [12,13], etc. In spite of these efforts, important questions remain to be answered as experimental data point toward complex thickness, angular, and temperature dependencies [14–18]. Although initial oversimplified theories attributed the dissipative damping-like component of the torque (even under time reversal) to spin Hall effect [19,20] and the reactive fieldlike component (odd under time reversal) to the inverse spin galvanic effect [6,21], this crude picture has been severely questioned by recent experiments. It remains unclear whether the torque components can be solely attributed to spin Hall effect, inverse spin galvanic effect, or a combination of both. In addition, a recent series of experiments [22–24] has identified unexpected torque components that are attributed to mechanisms beyond spin Hall and inverse spin galvanic effects [11–13]. A detailed discussion on these open questions can be found in Ref. [2].

To properly characterize and predict the behavior of spin-orbit torques in heterostructures, the model should be both comprehensive and transparent. As a matter of fact, such a model should ideally account for the realistic band structure of the heterostructure to treat bulk and interfacial spin-orbit effects on equal footing. But it should also be able to provide general trends that can serve as guidelines to experiments. To date, most models have addressed only certain aspects of the spin-orbit torques such as the interfacial inverse spin galvanic effect through model Hamiltonians [6–8,25–28] or spin Hall effect either through drift-diffusion or Boltzmann transport equation [20,29]. Whereas these approaches are quite transparent, their main limitation is their inability to treat both interface and bulk effects altogether and in particular the neglect of interfacial orbital hybridization that is known to be crucial in transition metal multilayers [30–32]. The common denominator between spin Hall and inverse spin galvanic effects is that they both stem from nonequilibrium orbital currents [33,34] or densities [35] that emerge from specific superposition of atomic orbitals. Consequently, the proper modeling of spin-orbit effects in heterostructures beyond the Rashba and spin Hall phenomenologies requires a multiorbital scheme.

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To date, the most accurate approach to compute multi-orbital transport properties is to rely on density-functional theory. This approach has been used extensively to compute spin and anomalous Hall effects in the bulk \cite{shekhar2020}, and has been recently extended to compute spin transport in heterostructures \cite{green2020}. Various techniques have been proposed including Wannier interpolation of the band structure \cite{manchon2020, kopp2020, korringa2020}, real-space Hamiltonians with tight-binding linear muffin-tin orbitals \cite{petersen2020, hedegaard2020}. While the first two methods are well adapted to compute the Kubo-Streda formula, the latter is suitable for two-terminal simulations, following the Landauer-Büttiker scheme. These different methods present the crucial advantage of accurately modeling the orbital hybridization across the whole structure. They are, however, computationally intensive, which makes them hardly adapted for systematic investigations (such as thickness dependence).

From this standpoint, developing a multiorbital tight-binding model is an interesting option as it reduces the size of the matrices to deal with numerically \cite{park2020, hedegaard2020}. When interfaced with density-functional theory, this method enables the accurate simulation of magnetic \cite{padovani2020} as well as transport properties \cite{petersen2020} in bulk materials. Using this approach, the intrinsic contribution to spin Hall effect has been computed in various materials (semiconductors \cite{green2020}, transition metals \cite{padovani2020, padovani2020, padovani2020}, topological insulators \cite{petersen2020}, Weyl semimetals \cite{park2020}, etc.) using the zero-temperature Berry curvature formula. Unfortunately, these results are only valid for vanishing disorder in the bulk and cannot be transposed to experimentally relevant setups where injection through interface dominates \cite{park2020}. To properly compute spin-charge conversion processes and spin-orbit torque, one needs to model the full heterostructure, including the interface \cite{park2020, manchon2020}. Recently, we applied this approach to a heterostructure made of a topological insulator capped with a ferro- or antiferro-) magnetic material \cite{green2020, green2020}. In this paper, each unit cell is modeled by a 4 × 4 Hamiltonian matrix regularized on a cubic lattice \cite{green2020}. Although quite crude, this approximation allowed us to model spin-orbit torque in various transport regimes and determine the minor role of spin Hall effect in these structures.

In the present paper, we use a multiorbital tight-binding model to compute the spin-orbit torque in transition-metal heterostructures. Whereas this method does not provide the accurate band structure obtained by density-functional theory, it retains the most prominent features of the density of states, atomic spin-orbit coupling, and interfacial orbital hybridization. It is also more flexible and computationally efficient, allowing for systematic characterization of the nonequilibrium properties of the heterostructure. In particular, we investigate the thickness and angular dependencies of the torque components and obtain a large “planar” dampinglike torque. We also investigate the nature of the self-torque, i.e., the spin-orbit torque taking place in the ferromagnet itself, and demonstrate that it can be substantial in spite of the large magnetic exchange \cite{park2020, manchon2020}.

II. MODEL AND FORMALISM

In this section, we first introduce a toy model to discuss how the interfacial orbital mixing gives rise to Rashba-like spin-orbit coupling. Then, we describe the tight-binding model of the heterostructure, and finally we expose the formalism we use to compute the transport properties.

A. Interfacial spin splitting with p and d orbitals

In centrosymmetric materials, such as the transition metals we consider in this paper, the spin Hall effect occurring in the bulk is usually attributed to intrinsic origin \cite{zhang2020, zhang2020}, i.e., to the Berry curvature of the wave functions. Following the scenario established by Tanaka et al. \cite{tanaka2020, tanaka2020}, Berry curvature in momentum space creates an orbital Hall current, which is spin polarized by turning on the atomic spin-orbit coupling. In contrast, little is known about the orbital origin of the interfacial Rashba spin-orbit coupling. Since the early works on this topic \cite{park2020, hedegaard2020}, it was proposed that upon inversion symmetry breaking, the spin-orbit coupling experienced by the Bloch electrons acquires a momentum-dependent Zeeman energy term, usually written

\begin{equation}
\mathcal{H}_R = \alpha_R \hat{\sigma} \cdot (\hat{\mathbf{p}} \times \mathbf{z}),
\end{equation}

where \( \alpha_R \) is called the Rashba parameter. Most early works on Rashba splitting have attributed its origin to the Zeeman splitting due to interfacial potential drop, \( \alpha_R = (\hbar/4mc^2) \int_{-\infty}^{\infty} \partial_z V \rho(z) dz \), where \( \partial_z V \) is the potential gradient and \( \rho(z) \) is the probability density distribution (see, e.g., Ref. \cite{park2020}). In fact, this contribution is an order of magnitude smaller than the Rashba splitting observed in \emph{ab initio} calculations \cite{park2020}. This discrepancy was addressed in a few recent works suggesting that the gradient of the Bloch-state wave function itself plays a central role in the onset of interfacial Rashba splitting \cite{petersen2020, hedegaard2020}. As a matter of fact, in their pioneering work, Petersen and Hedegård \cite{petersen2020} considered the Rashba spin splitting of Au (111) surface and proposed that the surface potential facilitates the superposition between \( p_z \) and \( p_x, p_y \) orbitals. This hybridization results in Rashba spin-orbit coupling when atomic spin-orbit coupling is turned on. A similar idea was put forward by several authors \cite{park2020, padovani2020, padovani2020}, suggesting that inversion symmetry breaking promotes the admixture between \( l \) and \( l \pm 1 \) orbitals. Recently, Park et al. \cite{park2020} suggested that the spin angular momentum texture in momentum space that characterizes Rashba spin-orbit coupling actually arises from a corresponding orbital angular momentum texture in the absence of spin-orbit coupling.

In this section, we wish to provide an explicit derivation of this effect and establish a direct connection between orbital mixture due to inversion symmetry breaking and Rashba-like spin-orbit coupling. Let us consider a chain of atoms, extended along \( x \) and with all three \( p_z, p_x, \) and \( p_y \) orbitals. We can discard the \( p_z \) orbitals from our discussion right away since they don’t couple to either \( p_x \), or \( p_y \). We now break the inversion symmetry by coupling this chain with another chain of atoms with only \( p_z \) orbitals. The system is depicted in Figs. 1 (a) and 1(b). The atoms of the bottom chain (with both \( p_z \) and \( p_x \)) are represented in gray and the atoms of the top chain (with \( p_z \) only) are in light blue. In the two-center tight-binding approximation and the \{\( p_x^\uparrow, p_y^\uparrow, p_z^\downarrow \)\} basis, the
Hamiltonian of this diatomic chain reads

$$\mathcal{H}_{\text{chain}} = \begin{pmatrix} \epsilon'_k & V_{zz} & V_{zx} \\ V_{zz} & \epsilon'_k & 0 \\ V_{zx} & 0 & \epsilon'_k \end{pmatrix}. \quad (2)$$

Here $p^\eta_k$ refers to the $\eta$th orbital of the bottom $(\eta = t)$ or bottom chain $(\eta = b)$, $V_{zz} = (V_0 + V_+ \cos k_a a/2)$ and $V_{zx} = -i(V_0 - V_+ \sin k_a a/2)$, $V_{b,t}$ being the Slater-Koster hopping integrals [75]. To keep our result analytically tractable, we assume that $\epsilon'_k = \epsilon''_k$. Then, we end up with three bands with dispersion,

$$\epsilon^0_k = \epsilon'_k, \quad \epsilon^±_k = \epsilon'_k \pm \frac{1}{2} \Delta_k, \quad (3)$$

with $\Delta_k = \sqrt{(\epsilon'_k - \epsilon''_k)^2 + 4t^2(V_{zz}^2 + V_{zx}^2)}$. The corresponding eigenstates read

$$|0\rangle = \frac{1}{\sqrt{|V_{zz}|^2 + |V_{zx}|^2}} (- V_{zx}|p^t_k\rangle + V_{zz}|p^b_k\rangle), \quad (4)$$

$$|+\rangle = \cos \chi |p^t_k\rangle + \frac{\sin \chi }{\sqrt{|V_{zz}|^2 + |V_{zx}|^2}} (V_{zx}|p^b_k\rangle + V_{zz}|p^t_k\rangle), \quad (5)$$

$$|-\rangle = - \sin \chi |p^b_k\rangle + \frac{\cos \chi }{\sqrt{|V_{zz}|^2 + |V_{zx}|^2}} (V_{zz}|p^b_k\rangle + V_{zx}|p^t_k\rangle), \quad (6)$$

where $\cos 2\chi = (\epsilon'_k - \epsilon''_k)/\Delta_k$. We now evaluate the orbital momentum on the bottom chain, and using $\langle p^\mu_k|L|p^\nu_k\rangle = i\gamma$, we get

$$\langle 0|L|0\rangle = -\frac{2\text{Im}[V_{zz}^* V_{zx}]}{|V_{zz}|^2 + |V_{zx}|^2}, \quad (7)$$

$$\langle +|L|+\rangle = \sin^2 \chi \frac{2\text{Im}[V_{zz}^* V_{zx}]}{|V_{zz}|^2 + |V_{zx}|^2}, \quad (8)$$

$$\langle -|L|-\rangle = \cos^2 \chi \frac{2\text{Im}[V_{zz}^* V_{zx}]}{|V_{zz}|^2 + |V_{zx}|^2}. \quad (9)$$

where $2\text{Im}[V_{zz}^* V_{zx}] = [(V_0)^2 - (V_+)^2] \sin k_a a$. This toy model shows that, due to the lack of inversion symmetry, the eigenstates of the diatomic chain acquire an orbital momentum that is odd in linear momentum $k$. This orbit-momentum locking results in orbital Edelstein effect [35], i.e., the electrical generation of an orbital magnetic moment.

When atomic spin-orbit coupling is turned on, the spin momentum of the Bloch electron aligns on its orbital momentum. Therefore, in the $\{|0\rangle, |+\rangle, |-\rangle\}$ basis, the spin-diagonal Hamiltonian of these Bloch states acquires an off-diagonal contribution,

$$\langle \xi_\mu \mathbf{L} \cdot \sigma \rangle = \mathcal{M}_{\xi_\mu} \frac{[(V_0)^2 - (V_+)^2] \sin k_a a}{\sqrt{(V_0)^2 + (V_+)^2 + 2V_0 V_+ \cos k_a a}} \delta_{\mu \gamma}, \quad (10)$$

where $\delta$ is the vector of Pauli spin matrices, and $\mathcal{M}_{\xi_\mu} = \text{diag}(-\xi_{\text{so}}^t, \xi_{\text{so}}^b \sin^2 \chi, \xi_{\text{so}}^b \cos^2 \chi)$, $\text{diag}(\ldots)$ being the diagonal matrix and $\xi_{\text{so}}^\eta$ the spin-orbit coupling energy of the $\eta$th chain $(\eta = t, b)$. This Hamiltonian explicitly connects the linear momentum $k_a$ with the spin momentum $\delta_\gamma$, resulting in Rashba and Dzyaloshinskii-Moriya effects [76]. This model can be straightforwardly extended to higher dimensions and higher order orbitals $(d, f, \text{etc.})$. In the case of a transition-metal interface, the orbital admixture required to obtain a spin density along $S_\gamma$ for an electron propagating along $x$ is typically $d_{z^2}$ or $d_{xy}$. A microscopic model of spin-orbit effects at interfaces should a minima contain these orbitals.

**B. Tight-binding model of the heterostructure**

We now move on to the description of the tight-binding model of our transition metal heterostructure, depicted in Fig. 2(a). This heterostructure consists of two adjacent metallic slabs with bcc crystal structure along the (001) direction, possessing the same lattice parameter. Each metallic slab consists of monolayers stacked on top of each other. Considering the ten $d$ orbitals, each monolayer adopts a square lattice described by the Hamiltonian

$$\mathcal{H}_{\text{mono}} = \begin{pmatrix} y_{xy}^k & 0 & t_{xy}^{k} \\ 0 & y_{yz}^k & t_{yz}^{k} \\ t_{xy}^{k*} & t_{yz}^{k*} & y_{xy}^k \end{pmatrix}, \quad (11)$$

where the parameters $y^k$ and $t^k$ are given explicitly in the Appendix. This Hamiltonian is written in the basis $(d_{xy}, d_{yz}, d_{zx}, d_{xz}, d_{x^2-y^2})$. This form is valid for each spin species, so the spin-dependent Hamiltonian reads $\mathcal{H}_{\text{mono}} \otimes \delta_\gamma$.
In addition, we define the exchange Hamiltonian, $H_{\text{ex}}$, as

$$H_{\text{ex}} = \frac{1}{2} \text{Diag}(\Delta_{xy}, \Delta_{yz}, \Delta_{xz}, \Delta_{\sigma^2 \rightarrow \pi^2}, \Delta_{\pi^2 \rightarrow \sigma^2}) \otimes \hat{\sigma} \cdot \mathbf{m}. \quad (12)$$

Here $\Delta_\nu$ is the exchange energy of the $\nu$th $d$ orbital and $\mathbf{m}$ is the ferromagnetic order parameter. Hence, the Hamiltonian for a square lattice monolayer is a $10 \times 10$ matrix. In addition, one needs to account for the spin-orbit-coupling matrix:

$$H_{\text{soc}} = \xi_{\text{soc}} \begin{pmatrix} 0 & i\hat{\sigma}_y & -i\hat{\sigma}_x & 0 & 2i\hat{\sigma}_z \\ -i\hat{\sigma}_y & 0 & i\hat{\sigma}_z & -i\sqrt{3}\hat{\sigma}_x & -i\hat{\sigma}_x \\ i\hat{\sigma}_x & -i\hat{\sigma}_z & 0 & i\sqrt{3}\hat{\sigma}_y & -i\hat{\sigma}_y \\ 0 & i\sqrt{3}\hat{\sigma}_x & -i\sqrt{3}\hat{\sigma}_y & 0 & 0 \\ -2i\hat{\sigma}_z & i\hat{\sigma}_z & i\hat{\sigma}_y & 0 & 0 \end{pmatrix}. \quad (13)$$

The Hamiltonian of a monolayer is therefore

$$H_0 = H_{\text{mono}} \otimes \hat{\sigma}_0 + H_{\text{ex}} + H_{\text{soc}}. \quad (14)$$

This monolayer is connected to the top nearest monolayer by the matrix $T_1$ whose elements are the nearest-neighbor hopping parameters $t_{\nu^\prime \nu}^ {k_{\nu^\prime}k_{\nu}}$ between orbital $d_{\nu}$ in the bottom layer and orbital $d_{\nu}$ in the top layer. The connection to the second top nearest monolayer is accounted for by the matrix $T_2$. The elements of both matrices are given explicitly in the Appendix. The Hamiltonian of one bcc slab is then defined

$$H_{\text{layer}} = \begin{pmatrix} H_0 & T_1 & 0 \\ T_1^\dagger & H_0 & T_2 \\ 0 & T_2^\dagger & H_0 \end{pmatrix}. \quad (15)$$

The matrix elements of $H_{\text{mono}}$, $T_1$, and $T_2$ are written in terms of the two-site Slater-Koster parameters (see Appendix) given for each slab in Table I. We adopt the parameters computed by Papaconstantopoulos [50] for bulk bcc Fe and bcc W. The lattice parameter of both slabs is set to that of bulk bcc W, $a_0 = 3.155$ Å, imposing $9\%$ lattice mismatch with bcc Fe whose bulk lattice parameter is 2.866 Å. Notice that the on-site energies of the ferromagnetic orbitals are rigidly shifted by an offset $\epsilon_0$ compared to their value in bulk Fe to allow for band-structure alignment between the ferromagnetic and nonmagnetic metals (see below). With these parameters, we determine the Hamiltonian for the nonmagnetic (NM) and ferromagnetic (FM) slabs, $H_{\text{NM}}$ and $H_{\text{FM}}$.

Finally, the heterostructure is obtained by stitching the two individual slabs together:

$$H = \begin{pmatrix} H_{\text{FM}} & T_{\text{FM}}^\dagger & H_{\text{NM}} \\ T_{\text{FM}} & H_{\text{FM}} & T_{\text{FM}}^\dagger \\ H_{\text{NM}} & T_{\text{NM}}^\dagger & H_{\text{NM}} \end{pmatrix}. \quad (16)$$

The hopping matrix $T_{\text{FM}}$ is simply given by $T_1$ and $T_2$ adopting the parameters of Table I. In the absence of further knowledge, the hopping parameters between the highest nonmagnetic layer and the lowest magnetic layer are taken as the average of the bulk hopping parameters of Fe and W. Ideally, one would need to fit the tight-binding parameters to the band structure of the heterostructure computed self-consistently from first principles [51], which remains out of the scope of the present paper but constitutes an appealing development of this work. Indeed, we emphasize that the nonmagnetic transition metal is expected to acquire interfacial magnetization by proximity with the ferromagnetic metal [31]. This induced magnetization is neglected in our model because our tight-binding parameters are that of the bulk materials. Nevertheless, a previous first-principles investigation of the Pt/Co(111) interface has shown that such an induced magnetization has a minor effect on the spin-orbit torque (see Fig. 7 in Ref. [40]).

Considering the numerous approximations we took (first- and second-nearest-neighbor hopping only, no self-consistent computation of the interfacial and exchange potentials, constrained lattice parameter, neglect of $s$ and $p$ orbitals, etc.), we do not expect our tight-binding model to accurately represent a realistic Fe/W bilayer. Nonetheless, we benchmarked our tight-binding model against the density of states of a Fe/W bilayer computed by density-functional theory to enforce its reliability. These simulations have been conducted using VIENNA AB INITIO SIMULATION PACKAGE (VASP) [77,78] with projector augmented wave (PAW) pseudo potential using the Generalized Gradient Approximation (GGA) with Perdew-Burke-Ernzerhof (PBE) exchange correlation [79,80]. The structure has been relaxed until forces on all the atoms go below 0.001 eV/Å allowing both the atomic coordinates and lattice vectors to change. We have used an energy cutoff of 500 eV. For self-consistent cycles, we used a $16 \times 16 \times 1$ k mesh and for the density of states, we used a $24 \times 24 \times 1$ k mesh. We neglected the effect of spin-orbit coupling and conducted a spin-polarized calculation, as we are interested in spin-resolved density of states. Including spin-orbit coupling does not make any drastic change in the total density of states.

The first-principles density of states of Fe(5)/W(7) projected on the $d$ orbitals only is reported in Fig. 3(a) together with the density of states obtained for our FM(5)/NM(7) system [Fig. 3(b)]. The figures in parenthesis indicate the number of monolayers and the density of states is defined $-\frac{1}{\pi} \text{Im}[\tilde{G}^R]$, where $\tilde{G}^R = (\epsilon - H + i\Gamma)$ is the retarded Green’s function and $\Gamma$ is the homogeneous broadening. The tight-binding density of states in Fig. 3(b) is obtained for a rigid energy shift $\epsilon_0 = 3.1$ eV, and the Fermi energy $E_F$ is fixed at 14 eV to qualitatively reproduce the balance between up and down Fermi electrons obtained by VASP. With these parameters, the total
number of electrons in the ferromagnetic and nonmagnetic metals are $n_{\text{FM}}^{\uparrow} = 4.6$, $n_{\text{FM}}^{\downarrow} = 2.34$, and $n_{\text{NM}}^{\uparrow} + n_{\text{NM}}^{\downarrow} = 4.73$.

We immediately observe a number of differences between the two densities of states in terms of bandwidth and peak position. These differences are attributed to the crude approximations of the tight-binding model mentioned above. Nevertheless, both densities of states display the same essential features: similar bandwidth, spin splitting of the ferromagnetic metal, large overlap between the two materials close to Fermi level, etc. Therefore, although our FM/NM heterostructure does not reproduce the ideal Fe/W case, it is a good representative of transition-metal heterostructures.

We conclude this discussion by considering the spin texture in momentum space. As explained above, symmetry breaking at the interface results in orbital texture in momentum space, which promotes the onset of spin-momentum locking in the presence of spin-orbit coupling \cite{74}. Figure 4 shows the band structure around the $\bar{\Gamma}$ point projected on the spin momentum components, $s_{x,y,z} = (\delta s_{x,y,z})$. In this calculation, the magnetization is set along $z$. Figure 4(a) displays the $s_x$ component when spanning the momentum between $\bar{Y}$ and $\bar{\Gamma}$ points, Fig. 4(b) displays the $s_y$ component when spanning the momentum between $\bar{X}$ and $\bar{\Gamma}$ points, and Fig. 4(c) displays the $s_z$ component along the $\bar{X} - \bar{\Gamma} - \bar{Y}$ path. The in-plane spin texture is antisymmetric in momentum and displays the $\sigma \sim z \times \mathbf{k}$ symmetry expected for Rashba spin-orbit coupling. In contrast, the $s_z$ component is symmetric and reflects the spin polarization of the bands due to magnetic exchange.

C. Transport formalism

The transport properties are computed using the Kubo-Streda formula \cite{41,81}. In this framework, the conductivity tensor reads

$$\sigma_{ij} = \frac{e^2 \hbar}{2\pi} \int d\varepsilon \partial \varepsilon f(\varepsilon) \text{Tr}[\hat{v}_j \hat{G}^R \hat{v}_i (\hat{G}^R - \hat{G}^A)].$$  \hspace{1cm} (17)

Here, \text{Tr} denotes the trace over the orbital, spin, and monolayer degrees of freedom and the sum over the Brillouin zone, $e = -|e|$ is the electron charge, and $\hat{v}_i = (1/\hbar) \partial_{\mathbf{k}} \mathcal{H}$ is the velocity operator. The local spin density on monolayer $\eta$ per unit electric field reads

$$S_\eta = \frac{e\hbar}{2\pi} \int d\varepsilon \partial \varepsilon f(\varepsilon) \text{Tr}[\hat{P}_\eta \otimes \sigma \hat{G}^R v_i (\hat{G}^R - \hat{G}^A)],$$  \hspace{1cm} (18)

where $\hat{P}_\eta$ is the projector on monolayer $\eta$. By construction, the matrix elements of $\hat{P}_\eta$ are equal to the $5 \times 5$ identity matrix $I_5$. 

![Fig. 4. Spin-resolved band structure for FM(5)/NM(7) bilayer: (a) $s_x$ along $\bar{Y}' - \bar{\Gamma} - \bar{Y}$, (b) $s_y$ along $\bar{X}' - \bar{\Gamma} - \bar{X}$, and (c) $s_z$ along $\bar{X} - \bar{\Gamma} - \bar{Y}$. Blue and red colors refer to opposite sign of the spin momentum.](174423-5)
at the position of layer $\eta$ and zero elsewhere:

$$\hat{\rho}_\eta = \begin{pmatrix} \cdots & 0 & \cdots \\ 0 & I_3 & 0 \\ \cdots & \cdots & \cdots \end{pmatrix}.$$  

(19)

The torque per unit electric field is defined as

$$T = \frac{e\hbar}{2\pi} \int d\varepsilon \delta_\varepsilon f(\varepsilon) \text{Tr}[\hat{\mathcal{H}} R \hat{\mathcal{V}} (\hat{\mathcal{G}}^R - \hat{\mathcal{G}}^A)],$$  

(20)

where $\tau = -m \times \partial_m \mathcal{H}$ is the torque operator. In the remaining of the paper, the conductivity of the slab is defined as $\sigma_{ij}/t$, $t$ being the thickness of the full heterostructure. The local spin density per unit electric field is in $\text{V}^{-1}\text{m}^{-1}$ and the torque, summed over the magnetic layer thickness, is expressed as a spin conductivity in the units of $(\hbar/2e)\Omega^{-1}$ $\text{m}^{-1}$. The disorder is accounted for through a homogeneous broadening $\Gamma$. Under this approximation, no higher order scattering events are taken into account (e.g., skew scattering, spin swapping, etc.).

III. CURRENT-DRIVEN SPIN-ORBIT TORQUES IN FM/NM HETEROSTRUCTURES

A. Spin-density profile

We first compute the current-driven spin-density profile throughout the heterostructure, when the magnetization $m$ points out of plane ($m[z]$). The two in-plane components, $S_x$ and $S_y$, are given in Figs. 5(a) and 5(b), respectively. The black curves correspond to the case where spin-orbit coupling is present in both ferromagnetic and nonmagnetic layers, while the red curves correspond to the case where only the ferromagnetic layer possesses spin-orbit coupling (see Sec. IV). When spin-orbit coupling is present in both ferromagnetic and nonmagnetic layers, we observe a clear accumulation of $S_x$ and $S_y$ components in the nonmagnetic metal. It is instructive to notice that the scale over which the spin density accumulates close to the interface is different for the two components. The $S_y$ component is localized close to the interface and vanishes quickly over about 10 MLs (ML, corresponding to about 1.3 nm), while $S_x$ slowly decays over a few tens of ML (i.e., about 5 nm). Notice also that $S_x$ penetrates deeper in the ferromagnetic layer than $S_y$. This distinction suggests that $S_x$ is controlled by nonlocal transport processes (e.g., scattering and diffusion), while $S_y$ is much more localized at the interface. Finally, a last important feature that distinguishes $S_x$ and $S_y$ is the presence of a nonvanishing $S_x$ component close to the outer surface of the nonmagnetic layer. These two features are consistent with the standard representation of spin-orbit torque as arising from diffusive spin Hall effect and interfacial Rashba-like effect. As a result, one expects the torque to display two components, conventionally referred to as fieldlike and dampinglike components, and reading

$$T_{FL} = \tau_{FL} m \times (z \times E),$$  

(21)

$$T_{DL} = \tau_{DL} m \times [(z \times E) \times m].$$  

(22)

B. Thickness dependence

We now address the thickness dependence of the two torque components, a property that has been investigated in numerous experiments [14,18,82–85]. To the best of our knowledge, such a thickness dependence has not been...
Figure 6. Dependence of the two torque components, (a) fieldlike torque and (b) dampinglike torque, as a function of the homogeneous broadening $\Gamma$, for different values of the transport energy, $E - E_F = 1$ eV (black), $E - E_F = 0$ eV (blue), and $E - E_F = -1$ eV (red). The inset displays the slab conductivity. Here the magnetization points perpendicular to the plane, along $z$.

The curves are calculated for a magnetization pointing along $z$ and for various disorder strengths, $\Gamma = 10$ meV (black), $\Gamma = 20$ meV (blue), $\Gamma = 50$ meV (red), and $\Gamma = 100$ meV (green).
interfacial orbital hybridization. This feature is therefore not general.

It is instructive to consider the thickness dependence of the torque efficiency, defined as the ratio between the torque and the conductivity of the heterostructure. This efficiency would be equivalent to the spin Hall angle if only spin Hall effect were present in the structure. The efficiency of the fieldlike and dampinglike torques is reported in Figs. 8(a) and 8(b), respectively, while the conductivity of the heterostructure is shown in the inset of (b), for reference. It is clear that the fieldlike torque efficiency is much larger for small thicknesses, as the current density is concentrated close to the interface. A similar feature is obtained for the dampinglike torque efficiency, defined as the ratio between the torque and the conductivity of the heterostructure. This efficiency would be equivalent to the spin Hall angle if only spin Hall effect were present in the structure. However, in the presence of additional torque components, the fieldlike torque efficiency is much smaller for small thicknesses, as the current density is concentrated close to the interface.

To complete this study, let us now consider the influence of the ferromagnetic layer thickness. Experimentally, it is found that the fieldlike component decreases strongly with the ferromagnetic layer thickness while the dampinglike component remains mostly constant [14]. We observe a similar feature in our calculations, shown in Figs. 7(c) and 7(d). The fieldlike component increases upon increasing the ferromagnetic layer thickness and saturates after about 10 MLs. The dampinglike component displays a similar increase as a function of the ferromagnetic layer thickness, but it also exhibits large quantum oscillations, which makes the systematic increase more difficult to see at first glance. This behavior is associated with the absorption of the transverse spin current by the ferromagnetic layer over the spin dephasing length. If the ferromagnetic layer thickness is thinner than the spin dephasing length, the injected spin current (or, equivalently, the spin density smearing into the ferromagnetic layer) is not entirely absorbed and is reflected back into the nonmagnetic layer, resulting in a reduced torque. Upon increasing the ferromagnetic layer thickness, more spin current is absorbed, resulting in an increase and saturation of the torque (see, e.g., Ref. [95]). This scenario was experimentally confirmed recently [96], but cannot be properly modeled using drift-diffusion theories due to the importance of quantum oscillations in this thickness range [20,87].

C. Angular dependence

The calculations presented above were all performed by setting the magnetization along $z$. Yet, several experimental [15,17,24] and theoretical studies [48,97–100] have pointed out that the spin-orbit torque does not reduce to the forms given in Eqs. (19)–(20). For the highest $C_{\infty}$ symmetry, Belashchenko et al. [48] proposed that the spin-orbit torque be written

$$
\mathbf{T} = -P^A_{\theta} \mathbf{m} \times (\mathbf{z} \times \mathbf{E}) + P^A_{\theta} (\mathbf{m} \cdot \mathbf{E}) \mathbf{m} \times (\mathbf{z} \times \mathbf{m})
- P^B_{\theta} \mathbf{m} \times [(\mathbf{z} \times \mathbf{E}) \times \mathbf{m}] + P^{\theta}_{\theta} (\mathbf{m} \cdot \mathbf{E}) \mathbf{m} \times \mathbf{z} + \ldots,
$$

(23)

where $P^N_{\theta} = \sum_n X_{2n} \mathcal{P}_{2n}(\cos \theta)$, $P_{2n}(x)$ being the Legendre polynomials. The first and third terms are simply the conventional fieldlike and dampinglike torques. The second and fourth terms can be referred to as “planar” fieldlike and “planar” dampinglike torques, respectively. These components are only nonzero when the magnetization lies along the applied electric field. In Ref. [98], these two planar components were obtained analytically and related to the presence of D’yakonov-Perel’ anisotropic spin relaxation. As a matter of fact, in such ultrathin magnetic heterostructures the spin component pointing perpendicular to the plane of the interface and that pointing in plane relax at different rates, which modifies the overall spin dynamics at the interface, resulting in these additional torque components.

To evaluate this angular anisotropy, we computed the two components $T_x$ and $T_y$ when varying the magnetization in the $(y, z)$ and $(z, x)$ planes. From Eq. (21), we expect

$$
T_x / \cos \theta = P^A_{\theta},
$$

(24)

$$
T_y / \cos \theta = P^B_{\theta},
$$

(25)

when the magnetization rotates in the $(y, z)$ plane, and

$$
T_x / \cos \theta = P^A_{\theta} - \sin^2 \theta P^N_{\theta},
$$

(26)

$$
T_y = P^B_{\theta} + \sin^2 \theta P^{\theta}_{\theta},
$$

(27)

where the magnetization rotates in the $(z, x)$ plane. By fitting these angular dependencies using Legendre polynomials, we obtain the first four components of the expansion of Eq. (21). These components are reported in Fig. 9 upon varying the thickness of the nonmagnetic metal.

The conventional fieldlike torque, reported in Fig. 9(a), dominates all the other components and exhibits almost no...
contributes to the torques reported in Fig. 9.

small (see Sec.IV) and it is unlikely that such a mechanism
in our case, the spin-orbit coupling of the ferromagnet remains quite
planar Hall effect from the bulk of the ferromagnet. In our
complete this discussion, we emphasize that Safranski
the conventional (spin Hall-driven) dampinglike torque. To
torque is associated with bulk mechanisms and accompanies
facial (Rashba) fieldlike torque, while the planar dampinglike
and can be seen at the companion of the conventional inter-
progressive saturation is similar to the one expected for spin Hall-driven damping torque originating
of the nonmagnetic layer, as mentioned in Sec. III B.
Therefore, the results reported in Fig. 9 suggest that the
planar fieldlike torque is associated with an interfacial effect
and can be seen at the companion of the conventional inter-
facial (Rashba) fieldlike torque, while the planar dampinglike
torque is associated with bulk mechanisms and accompanies
the conventional (spin Hall-driven) dampinglike torque. To
complete this discussion, we emphasize that Safranski et al.
reported a planar Hall torque that they attributed to the
planar damping-like torque, reported in Fig. 9(b), is about one order
of magnitude smaller, exhibits the sign reversal discussed
previously, and displays a sizable angular dependence at small
thicknesses ($B_2 \gg B_{4,6}$). This angular dependence vanishes
upon increasing the thickness of the nonmagnetic layer.
Interestingly, the two planar components exhibit a radically
different behavior. First, both components are comparable in
magnitude with the dampinglike torque, which means that
they play a crucial role in current-driven dynamics and cannot
be neglected. Second, the planar fieldlike torque [Fig. 9(c)]
exhibits a substantial angular dependence ($A'_0 \approx A'_2 \gg A_{4,6}$)
that saturates after only a few monolayers. This indicates that
this component is mostly of interfacial origin, in agreement
with the D’yakonov-Perel’ scenario evoked in Ref. [98].
Finally, the planar dampinglike torque presents a surprising
behavior [Fig. 9(d)]. It displays almost no angular dependence
(except at small thicknesses), and increases steadily over a few tens of monolayers before reaching saturation at large
thicknesses. This progressive saturation is similar to the one
expected for spin Hall-driven damping torque originating
from the nonmagnetic layer, as mentioned in Sec. III B.

IV. SELF-TORQUE IN THE FERROMAGNET

To complete this study, we now turn off the spin-orbit cou-
pling of the nonmagnetic layer. The spatial profile of the spin
density is shown in Fig. 5, red curves. The features described
above survive: $S_z$ is more delocalized than $S_y$, although their
magnitude is much (three or four times) weaker than in the case
where spin-orbit coupling is present in both layers. The
thickness dependence is shown in Fig. 10, blue curves.
The black curves represent the case where the spin-orbit
coupling is present in both layers and serves as a reference.
The fieldlike torque starts slightly positive [Fig. 10(a)],
switches sign around about 10 MLs, and increases negatively
until reaching saturation at about 50 MLs. The damping-
like torque shows a similar behavior. It also starts slightly
positive [Fig. 10(b)], switches sign about 15 MLs, and in-
creases negatively until reaching saturation at about 60 MLs.
It is interesting to note that the self-fieldlike torque has
the same sign as the case where spin-orbit coupling is present
everywhere, whereas the self-dampinglike torque is opposite.

To understand this distinct behavior, we compute the de-
pendence of the anomalous Hall conductivity and torque
components as a function of the spin-orbit coupling energy
of the individual layers. The results are reported in Fig. 11.
The black lines correspond to the case where the spin-orbit
coupling is in the nonmagnetic metal only, whereas the blue
lines correspond to the case where the spin-orbit coupling
is in the ferromagnet only. The anomalous Hall conductivity
[Fig. 11(a)] and dampinglike torque [Fig. 11(c)] both change
sign depending on which layer possesses spin-orbit coupling.
In contrast, the fieldlike torque remains negative, irrespective
of where the spin-orbit coupling is [Fig. 11(b)].

The fieldlike torque, as explained in Sec. II A, is asso-
ciated with the interfacial Rashba-like spin-orbit coupling
whose sign is governed by the interfacial orbital hybridization.
Therefore, for a given spin-orbit coupling strength, its sign
FIG. 11. Dependence of the (a) anomalous Hall conductivity, (b) fieldlike, and (c) dampinglike components of the spin-orbit torque upon varying the spin-orbit coupling $\xi_{so}$. The black lines represent the case where the spin-orbit coupling of the ferromagnet is set to zero, whereas the blue lines represent the case where the spin-orbit coupling of the nonmagnetic metal is set to zero. In this calculation, we set the nonmagnetic metal thickness to 40 MLs and the ferromagnet thickness to 7 ML. The curves are calculated for a magnetization pointing along $z$ and for $\Gamma = 50$ meV.

is opposite on the two sides of the interface [see Eq. (9)]. This seems contradictory with the results of Fig. 11(b) and suggests that the sign of the spin-orbit coupling experienced by the Bloch states of the nonmagnetic layer is opposite to the one experienced by the Bloch states of the ferromagnet. This observation is consistent with Hund’s third rule that states that for materials with more-than-half-filled electronic shells such as Fe, the spin and orbital momenta are aligned with each other, while for materials with less-than-half-filled electronic shells like W, they are antialigned. Since our tight-binding model is parameterized on these two elements, it is reasonable that Hund’s third rule applies. As a consequence, the opposite potential drop felt by Bloch states on each side of the interface is compensated by the opposite effective spin-orbit coupling, and the fieldlike torque is the same whether the spin-orbit coupling is on the ferromagnet or on the nonmagnetic layer.

In contrast, the dampinglike torque at large thicknesses and the anomalous Hall conductivity are not associated with interfacial potential drop, but rather with the (spin) Berry curvature of the bulk material. It is therefore solely governed by the effective spin-orbit coupling experienced by the Bloch electrons and is opposite when switching the spin-orbit coupling from the ferromagnet to the nonmagnetic layer.

The dependence as a function of the ferromagnetic layer thickness is reported in Figs. 10(c) and 10(d) for the fieldlike and dampinglike torques, respectively. We obtain similar thickness dependence as in the case where spin-orbit coupling is present in both layers, reflecting the importance of the spin dephasing length. Using our realistic parameters, the self-torque we obtain is about four to five times smaller in magnitude compared to the torque arising from the nonmagnetic metal, consistent with the relative magnitude of the spin-orbit coupling (about 65 meV in Fe compared to 360 meV in W). These calculations support an idea that was put forward in Ref. [59]: The spin Hall current generated inside the ferromagnetic layer can create an efficient torque on the magnetic order as long as the two opposite interfaces are dissimilar.

V. CONCLUSION

Using a multiorbital tight-binding model, we computed the spin-orbit torque in a transition metal heterostructure, treating bulk and interfacial spin-orbit effects coherently and on equal footing. Thickness and angular dependencies of the torque show that it possesses four sizable components, the conventional fieldlike and dampinglike torques, as well as two planar components that vanish when the magnetization lies out of plane. The conventional fieldlike torque is entirely controlled by the interface, as expected from interfacial inverse spin galvanic effect, while the dampinglike torque possesses two components, an interfacial one dominating at small thicknesses and a bulk contribution dominating at large thicknesses. The former is attributed to the intrinsic interfacial Berry-curvature-driven damping torque [94], whereas the latter is associated with the spin Hall effect generated in the bulk of the nonmagnetic metal.

Interestingly, the planar fieldlike torque shows substantial angular dependence and is of interfacial origin, like the conventional fieldlike torque. In contrast, the planar dampinglike torque does not exhibit angular dependence and increases with the nonmagnetic metal thickness, indicating that it originates from the bulk of the nonmagnetic layer, similar to the conventional spin Hall-driven damping torque. Our results demonstrate that these four torque components are present in any transition-metal heterostructures and must be taken into account when interpreting the experimental data, and in particular the current-driven magnetization dynamics.

Finally, we investigate the self-torque exerted on the ferromagnet when spin-orbit coupling of the nonmagnetic metal is turned off. Our results suggest that the spin accumulation that builds up inside the ferromagnet can be large enough to induce magnetization excitations.

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APPENDIX: HOPPING INTEGRALS

The parameters appearing in Eq. (10) are

\[
\begin{align*}
\gamma_{xy}^{k} &= \varepsilon_{xy} + (3V_{x}^{2} + V_{y}^{2}) \cos \tilde{k}_{x} \cos \tilde{k}_{y}, \\
\gamma_{yz}^{k} &= \varepsilon_{yz} + 2(V_{x}^{2} + V_{z}^{2}) \cos \tilde{k}_{x} \cos \tilde{k}_{z}, \\
\gamma_{zx}^{k} &= \varepsilon_{zx} + 2(V_{y}^{2} + V_{z}^{2}) \cos \tilde{k}_{y} \cos \tilde{k}_{z}, \\
\gamma_{xy}^{k,z} &= \varepsilon_{xy} + (V_{x}^{2} + 3V_{y}^{2}) \cos \tilde{k}_{x} \cos \tilde{k}_{y}, \\
\gamma_{yz}^{k,x} &= 2(V_{x}^{2} - V_{y}^{2}) \cos \tilde{k}_{x} \sin \tilde{k}_{y}, \\
\gamma_{zx}^{k,y} &= \sqrt{3}(V_{x}^{2} - V_{z}^{2}) \sin \tilde{k}_{x} \sin \tilde{k}_{y}, \\
\end{align*}
\]

where \( V_{\sigma,\pi,\delta} \) are the two-site second-nearest-neighbor hopping integrals in the \((x, y)\) plane [75], as depicted in Fig. 2(b), and \( \tilde{k}_{xy} = k_{xy}a_{0}/2 \). The matrix elements of the nearest-neighbor hopping matrix \( \mathbf{T}_{1} \) are

\[
\begin{align*}
t_{xy}^{k,x} &= (V_{x}^{2} + V_{y}^{2}) \cos \tilde{k}_{x} + \cos \tilde{k}_{y}, \\
t_{yz}^{k,y} &= (V_{y}^{2} + V_{z}^{2}) \cos \tilde{k}_{y} + 2(V_{x}^{2} + V_{y}^{2}) \cos \tilde{k}_{x}, \\
t_{zx}^{k,z} &= (V_{z}^{2} + V_{x}^{2}) \cos \tilde{k}_{z} + 2(V_{y}^{2} + V_{z}^{2}) \cos \tilde{k}_{y}, \\
t_{xz}^{k,x} &= \frac{1}{8}(3V_{x}^{4} + 4V_{y}^{4} + 9V_{z}^{4}) \cos \tilde{k}_{x} + \cos \tilde{k}_{y}, \\
t_{zy}^{k,y} &= \frac{1}{8}(V_{y}^{4} + 12V_{z}^{4} + 3V_{x}^{4}) \cos \tilde{k}_{y} + \cos \tilde{k}_{z}, \\
\end{align*}
\]

where we used the shorthand notation \( t_{0}^{k} = t_{0}^{k} \), and

\[
\begin{align*}
t_{xy}^{k} &= -i(V_{x} - V_{y}) \sin \tilde{k}_{y}, \\
t_{yz}^{k} &= -i(V_{y} - V_{z}) \sin \tilde{k}_{x}, \\
t_{zx}^{k} &= 3i(V_{z} - V_{x}) \sin \tilde{k}_{y}, \\
t_{xz}^{k} &= -\frac{3i}{4}(V_{z} - V_{x}) \sin \tilde{k}_{x}, \\
t_{zy}^{k} &= 0. \\
\end{align*}
\]

Again, \( V_{\sigma,\pi,\delta} \) are the two-site second-nearest-neighbor hopping integrals [75], as depicted in Fig. 2(b). The second-nearest-neighbor hopping matrix \( \mathbf{T}_{2} \) reads

\[
\mathbf{T}_{2} = \text{Diag}(V_{x}^{2}, V_{y}^{2}, V_{z}^{2}, V_{x}^{2}, V_{y}^{2}, V_{z}^{2}),
\]

which connects the \( n \)th monolayer to the \( n+2 \)th monolayer.

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