Spin-orbit coupling effects on spin-dependent inelastic electronic lifetimes in ferromagnets

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For the 3d ferromagnets iron, cobalt, and nickel we compute the spin-dependent inelastic electronic lifetimes due to carrier-carrier Coulomb interactions including spin-orbit coupling. We find that the spin-dependent density of states at the Fermi energy does not, in general, determine the spin dependence of the lifetimes because of the effective spin-flip transitions allowed by the spin mixing. The majority and minority electron lifetimes computed including spin-orbit coupling for these three 3d ferromagnets do not differ by more than a factor of 2, and agree with experimental results.

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The theoretical and experimental characterization of spin dynamics in ferromagnetic materials due to the interaction with short optical pulses has become an important part of research in magnetism [1–6]. In this connection, spin-dependent hot-electron transport processes in metallic heterostructures have generated enormous interest in the past few years [7]. In particular, superdiffusive-transport theory has played an increasingly important role in the quantitative interpretation of experimental results [4,6,8]. Superdiffusive-transport theory, which was introduced and comprehensively described in Refs. [9,10], uses spin- and energy-dependent electron lifetimes as the input [10], and its quantitative results for hot-electron transport on ultrashort time scales in ferromagnetic materials rely heavily, to the best of our knowledge, on the relation between majority and minority electrons for these materials.

The spin-dependent lifetimes that are used for hot-electron transport, both in ferromagnets and normal metals, are the so-called “inelastic lifetimes.” These state- (or energy-) dependent lifetimes result from outscattering processes due to the Coulomb interaction between an excited electron and the inhomogeneous electron gas in the system. The lifetimes can be measured by tracking optically excited electrons using spin- and time-resolved two-photon photoemission (2PPE) [11,12] and can be calculated as the broadening of the electronic spectral function using many-body Green’s function techniques [13,14]. The problem of the accurate determination of inelastic lifetimes has fueled method development on the experimental and theoretical side [15], but has always suffered from the presence of interactions (electron-phonon, surface effects) that cannot be clearly identified in experiment and are difficult to include in calculations. Qualitative agreement was reached for the spin-integrated lifetimes in simple metals and iron [16]. However, even advanced quasiparticle calculations, including many-body $T$-matrix contributions, have yielded a ratio between majority and minority lifetimes which is in qualitative disagreement with experiment for some ferromagnets. Largely due to this discrepancy, a consensus has yet to be reached on the importance of spin-dependent hot-electron transport processes because, in numerical studies on nickel [4,6,10], the theoretical ratio of about 8 [16] has been used, while the experimental result [17] is 2. The problem is exacerbated by recent experimental results that point toward a similar disagreement for cobalt [12].

In light normal metals and ferromagnets spin-orbit coupling generally leads to very small corrections to the single-particle energies, i.e., the band structure, but it changes the single-particle states qualitatively by introducing a state-dependent spin mixing. With spin-orbit coupling, the average spin of an electron can be changed due to electron-phonon and electron-impurity interactions [18–21]. This is also true for the two-particle Coulomb interaction [22,23], as long as one monitors only the average spin of one of the scattering particles, as is done in lifetime measurements by two-photon photoemission experiments. The spin mixing in the wave functions due to spin-orbit coupling has not been included in existing lifetime calculations for 3d ferromagnets, aluminum, and lead [16,24–26]. However, recently there have been developments to include spin-orbit coupling in many-body perturbation theory [27,28].

This Rapid Communication presents results for electron lifetimes in metals and spin-dependent lifetimes in ferromagnets that include spin-orbit coupling in the wave functions and the band structure. We show that spin-orbit coupling, in particular, the spin mixing in the wave functions, is important for electron lifetimes in metals, in general. The ratio between the majority and minority lifetimes calculated with spin mixing is now in agreement with experiment [11,12,17]. We believe that our calculated electronic lifetimes should be used as an accurate input for calculations of spin-dependent hot-electron dynamics in ferromagnets.

We first discuss briefly our theoretical approach to calculating the lifetimes. We start from the dynamical and wavevector-dependent dielectric function $\varepsilon(\mathbf{q},\omega)$ in the random phase approximation (RPA) [13,14,25,26]. Our approach (cf. Ref. [29]) evaluates the wave-vector summations in $\varepsilon(\mathbf{q},\omega)$ without introducing an additional broadening of the energy-conserving $\delta$ function. This procedure removes a parameter whose influence on the calculation for small $q$ is not easy to control and which would otherwise need to be separately tested over the whole energy range.

The $\mathbf{k}$- and band-resolved electronic scattering rates, i.e., the inverse lifetimes, $\gamma_k^\nu = (\tau_k^\nu)^{-1}$, are calculated using the expression [25,26]

$$
\gamma_k^\nu = \frac{2}{\hbar} \sum_{\mathbf{q},\mu} \frac{\Delta q^3}{(2\pi)^3} V_q |B^{\nu\mu}_{\mathbf{k}+\mathbf{q}}|^2 \frac{f^\mu_{\mathbf{k}+\mathbf{q}}}{\varepsilon(\mathbf{q},\Delta E)} \left| \frac{\Im \varepsilon(\mathbf{q},\Delta E)}{\varepsilon(\mathbf{q},\Delta E)^2} \right|^2.
$$

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Here, the band indices are denoted by $\mu$ and $\nu$, and $\vec{k}$ and $\vec{q}$ denote wave vectors in the first Brillouin zone (1BZ). The energies $\epsilon_k^\mu$, occupation numbers $f_k^\mu$, and overlap matrix elements $B_{kq}^{\mu\nu} = \langle \psi_k^\mu | e^{i\vec{q}\cdot\vec{r}} | \psi_k^\nu \rangle$ are extracted from the ELK density functional theory (DFT) code [30], which employs a full-potential linearized augmented plane wave (FP-LAPW) basis. Last, $V_q = e^2/(\epsilon_0 q^2)$ denotes the Fourier transformed Coulomb potential and $\Delta E = \epsilon_k^\mu - \epsilon_k^\nu$ is the energy difference between the initial and final state. For negative $\Delta E$, the distribution function has to be replaced by $-1 - (1 - f_{k+\vec{q}}^\mu)$. By using the overlap matrix elements as defined above we neglect corrections due to local field effects. In the language of many-body Green’s functions, this corresponds to an on-shell $G_0 W_0$ calculation [16,26], where the screened Coulomb interaction ($W_0$) is obtained from the full RPA dielectric function. The $\vec{k}$- and band-dependent wave functions that result from the DFT calculations including spin-orbit coupling are of the form $| \psi_{\vec{k}}^{\mu} \rangle = a_{\vec{k}}^{\mu\uparrow} | \uparrow \rangle + b_{\vec{k}}^{\mu\downarrow} | \downarrow \rangle$ [18], where $| \sigma \rangle$ are spinors identified by the spin projection $\sigma = \uparrow, \downarrow$ along the magnetization direction, which is [001] for Fe(bcc), Ni(fcc), and [0001] for Co(hcp). According to whether $| a_{\vec{k}}^{\mu\uparrow} |^2$ or $| b_{\vec{k}}^{\mu\downarrow} |^2$ is larger, we relabel each eigenstate by its dominant spin contribution $\sigma$, so that we obtain spin-dependent lifetimes $\tau_{\vec{k}}^\sigma$. Our choice of quantization axis is such that $\sigma = \uparrow$ denotes majority carrier states and $\sigma = \downarrow$ minority carrier states. Due to the existence of several bands (partially with different symmetries) in the energy range of interest and the anisotropy of the DFT bands $\epsilon^{(\nu)}(\vec{k})$, several lifetimes $\tau_{\vec{k}}^\sigma$ can be associated with the same spin and energy. When we plot these spin- and energy-dependent lifetimes $\tau(\vec{k})$ in the following, in particular, Figs. 1 and 2, this leads to a scatter of $\tau(\vec{k})$ values.

Figures 1 and 2 display the calculated energy- and spin-resolved lifetimes $\tau(\vec{k})$ around the Fermi energy for cobalt and nickel. The spread of lifetimes at the same energy, which was mentioned above, can serve as an indication for the possible range of results for measurements of energy-resolved lifetimes. These “raw data” are important for the interpretation of the theoretical results because they already show two important points. First, we checked that there is no good Fermi-liquid type fit to these lifetimes. Second, even if one fits the lifetimes in a restricted energy range by a smooth $\tau(E)$ curve, this ignores the spread of lifetimes, which can be quite sizable, as shown in Figs. 1 and 2. We believe that such a spread of electronic lifetimes, in particular, in the range around 1 eV above the Fermi energy, should be important for the interpretation of photoemission experiments in this energy range, and when these results are used as the input in hot-electron transport calculations.

Figure 1 shows the energy- and spin-resolved lifetimes in cobalt. In addition to the longer lifetimes close to the Fermi energy, hole lifetimes in excess of 5 fs occur at the top of some $d$ bands around $-1.5, -1.2,$ and $-1$ eV. For electronic states with energies above 0.5 eV, longer lifetimes occur at some $\vec{k}$ points. There are also $k$ states with a pronounced spin asymmetry in the lifetimes (see the discussion below). Another important property of cobalt is the existence of two different conduction bands, which intersect the Fermi surface with different slope. This leads to two rather well-defined lifetime curves, both for electrons and holes. This can be best seen between $-0.6$ and $0$ eV, where the two curves are shifted by about 0.2 eV.

The calculated lifetimes in nickel (see Fig. 2) do not show a pronounced influence of $d$ bands and/or anisotropy below the Fermi energy as in cobalt, which is due to the smaller number of bands in the vicinity of the Fermi energy. However, there is a clear spin dependence of electronic lifetimes, which is most pronounced around 0.4 eV, but persists almost up to 2 eV.

In the following, we will mainly be concerned with lifetimes above 0.3 eV above the Fermi energy, which is the interesting energy range for the interpretation of photoemission experiments and hot-electron transport calculations, because close to the Fermi energy the influence of phonons is expected to become more pronounced and can lead to significantly shorter lifetimes than those predicted by a calculation that includes only the Coulomb interaction. To facilitate comparison with experiment, we average the lifetimes in each spin channel in bins of 100 meV and denote the result by $\bar{\tau}(\vec{k})$. The standard deviation of the averaging process then yields “error bars” on
for larger energies, a trend that agrees extremely well with measurements [11,12,17]. To put this result into perspective we note that the experimental data in Ref. [17] were compared with a theoretical model based on the random-\(k\) approximation [33]. If the random-\(k\) interaction matrix elements are taken to be spin and energy independent, the majority and minority relaxation times are determined by double convolutions over the spin-dependent density of states (DOS) [16]. It was found that the experimental results were not in agreement with the ratio of the DOS at the Fermi energy, which led the authors of Ref. [17] to speculate that the matrix elements for parallel and antiparallel spins should be different due to the Pauli exclusion principle. In our calculations, the effective spin dependence of the matrix elements is caused exclusively by the spin mixing due to spin-orbit coupling, but the effect is the same: It makes the ratio of the lifetimes different from the spin-dependent DOS at the Fermi energy.

In Fig. 3(c) we turn to nickel. Here, as in the case of iron, the average lifetimes are slightly larger than the measured ones [17] (not shown), but due to the small anisotropy in the band structure, the lifetimes in nickel show the smallest error bars and thus an extremely well-defined spin dependence. Only our calculated majority electron lifetimes are similar to earlier \textit{ab initio} evaluations [15,16,31], but there is an important discrepancy in the ratio \(\tau_\uparrow/\tau_\downarrow\): The inset of Fig. 3(c) shows

\begin{figure}[h]
\centering
\includegraphics[width=\textwidth]{fig3}
\caption{(Color online) Energetically averaged majority (blue up-triangles) and minority (red down-triangles) lifetimes for (a) Fe, (b) Co, and (c) Ni. The error bars denote the standard deviation obtained from the scatter of the lifetimes as shown in Figs. 1 and 2. The insets show the calculated ratio of majority and minority electrons \((\circ), \tau_\uparrow/\tau_\downarrow\), in comparison with experimental data, where the \(\bullet\ (\times)\) correspond to values extracted from Refs. [11,17] (Ref. [12]).}
\end{figure}

the \(\bar{\tau}(E)\) values. Note that this procedure does not correspond to a “random-\(k\)” approximation.

Figure 3 displays the averaged electron lifetimes determined from the data shown in Figs. 1 and 2. As insets we have included the ratio of majority and minority lifetimes, \(\tau_\uparrow/\tau_\downarrow\), together with experimental data [11,12,17] for iron, cobalt, and nickel. Figure 3(a) shows that there is only a very weak spin dependence for iron, and the agreement of the ratio \(\tau_\uparrow/\tau_\downarrow\) with experiment [17] and recent investigations [15,16,31] is quite good, but there is a slight disagreement with earlier, semiempirical studies [32]. However, even an increase of the ratio around 0.5 eV in the experiment [17] is well reproduced in our results.

The averaged lifetimes of cobalt, which are shown in Fig. 3(b), agree quite well with the experimental lifetimes [11,12], but the large error bars extend to a much wider energy range than in iron. This can be traced back to the scatter of lifetimes in Fig. 1. The corresponding figure for iron (not shown) exhibits a much smaller scatter. The ratio of majority and minority electron lifetimes [see the inset in Fig. 3(b)] is around 1 below 0.5 eV and increases to \(\tau_\uparrow/\tau_\downarrow \simeq 2\)

\begin{figure}[h]
\centering
\includegraphics[width=\textwidth]{fig4}
\caption{(Color online) Calculated energy-resolved electronic lifetimes for cobalt and nickel in analogy to Fig. 3 without the inclusion of spin-orbit coupling effects.}
\end{figure}
a ratio of $\tau_\uparrow/\tau_\downarrow \approx 2$, which is independent of energy above 0.4 eV. This result compares extremely well with experiment, and should be contrasted with the calculated result of Ref. [16] for $\tau_\uparrow/\tau_\downarrow \approx 8$ around 0.5 eV. These GW calculations (even with a $T^\ast$-matrix approach) gave very similar results to those of the random-$k$ approximation [16] in the energy range above 0.5 eV. This indicates that the resulting spin asymmetry $\tau_\uparrow/\tau_\downarrow \approx 6–8$ is solely determined by the spin-dependent DOS $D_{\sigma}(E)$. Indeed, one has $D_{\uparrow}(E_F)/D_{\downarrow}(E_F) \approx 8$. With the inclusion of spin-orbit coupling, which gives rise to effective spin-flip transitions, the spin asymmetry is no longer determined by the spin-dependent DOS alone. Without knowledge of the microscopic mechanism, the effect of a strongly enhanced spin-flip matrix element had already been introduced by hand in Ref. [17], and was recognized to improve the agreement between a random-$k$ calculation and experiment. We would like to briefly compare the results of Fig. 3, which generally shows relatively small and weakly energy-dependent spin asymmetry of the lifetimes in 3d-ferromagnets, with those for Heusler compounds [34]. In the latter materials we have found a more pronounced energy dependence of the spin asymmetry as well spin asymmetries around 5, which are generally realised at the bottom of minority bands.

To conclude the discussion of the ferromagnets, we comment on the spin-integrated lifetimes which can be obtained from the spin-dependent lifetimes, but are not shown here. Compared with the experimental lifetimes of Ref. [17], we generally find an agreement for energies above 0.5 eV that is on par with earlier calculations [15–17,31]. For energies below 0.5 eV where the error bars on the averaged lifetimes are largest, the calculated lifetimes are larger than the measured ones, but in this energy range a good agreement with experiments cannot be expected because of scattering processes, which appear as elastic due to the energy resolution of the photoemission experiments.

To underscore the importance of spin-orbit coupling for lifetime calculations, we also briefly show calculated results for electronic lifetimes in cobalt and nickel without the inclusion of spin-orbit coupling effects in Fig. 4. In this case, which essentially corresponds to the GW approximation of Ref. [16], the minority lifetimes calculated remain nearly unchanged in comparison to Fig. 3, whereas the majority electron lifetimes are changed drastically. As further displayed in Fig. 4, this results in a pronounced spin asymmetry. In particular, the ratio between majority and minority electron lifetimes increases from 4 (3) at 1.2 eV to 11 (6) at 0.2 eV in cobalt (nickel).

In conclusion, we presented ab initio results for spin-dependent electronic lifetimes in ferromagnets including spin-orbit coupling in the wave functions and energies. We found that the electronic lifetimes in iron exhibit no visible spin dependence in the range of $\pm 2$ up to 3 eV, in agreement with earlier results, whereas the ratio $\tau_\uparrow/\tau_\downarrow$ between majority and minority lifetimes does not exceed 2 for cobalt and nickel. Our results agree well with experimental data, but differ from earlier calculations, which found that $\tau_\uparrow/\tau_\downarrow$ was essentially determined by the spin-dependent density of states. We showed that, by allowing for effectively spin-changing transitions as contributions to the lifetime, spin-orbit coupling is the essential ingredient that can make the spin asymmetry of the electronic lifetimes much smaller than the spin asymmetry of the density of states. Inclusion of our calculated spin-dependent lifetimes in transport calculations should make it possible to more accurately characterize the influence of spin-dependent hot-electron transport on magnetization dynamics.

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