Elliott-Yafet Spin-Phonon Relaxation Times from First Principles

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We present a first-principles approach for computing the phonon-limited $T_1$ spin relaxation time due to the Elliott-Yafet (EY) mechanism. Our scheme combines fully-relativistic spin-flip electron-phonon interactions with an approach to compute the effective spin of band electrons in materials with inversion symmetry. We apply our method to silicon and diamond, for which we compute the temperature dependence of the spin relaxation times and analyze the contributions to spin relaxation from different phonons and valley processes. The computed spin relaxation times in silicon are in excellent agreement with experiment in the 50–300 K temperature range. In diamond, we predict intrinsic spin relaxation times of $540 \mu s$ at 77 K and 2.3 $\mu s$ at 300 K. Our work enables precise predictions of spin-phonon relaxation times in a wide range of materials, providing microscopic insight into spin relaxation and guiding the development of spin-based quantum technologies.

Introduction. Spin relaxation in centrosymmetric crystals primarily occurs through the Elliott-Yafet (EY) mechanism [1, 2], in which spin decoherence can be mediated by electron-phonon (e-ph) or electron-defect interactions. Phonons typically dominate EY spin relaxation near room temperature, and often limit the performance of spin-based devices in spintronics [3–5] and quantum technologies [6–9]. Recent advances are pushing spin manipulation to new frontiers [10–13], so understanding in detail how electron spins interact with phonons is important for both technological and fundamental reasons.

Accurately predicting spin-phonon relaxation processes remains an open problem, particularly due to the challenge of quantifying spin-flip e-ph interactions [2]. Calculations of EY spin relaxation have mainly relied on empirical models [14, 15] and symmetry analysis [16–19], yet these approaches are laborious even for simple materials and not geared toward quantitative predictions. Attempts have also been made to study spin relaxation from first principles by assuming a direct proportionality between spin-flip and momentum-scattering e-ph interactions [20], or between spin-flip and momentum relaxation times [21]. However, these assumptions hold only for simple model potentials [22–24] as the spin-flip and momentum-scattering processes can differ greatly depending on the electronic wave function, spin texture and phonon perturbation [19, 25].

Recently developed first-principles methods for computing e-ph interactions and relaxation times [26] are promising for studying EY spin-phonon relaxation. Their typical workflow [27] involves density functional theory (DFT) calculations of the ground state and electronic band structure, combined with density functional perturbation theory (DFPT) [28] to compute the phonon dispersions and e-ph perturbation potentials, followed by interpolation of the e-ph coupling matrix elements to fine Brillouin zone (BZ) grids. However, this workflow cannot be applied as is to investigate spin-flip e-ph interactions because the spin information is lost when one computes the e-ph matrix elements. For example, the electronic states in centrosymmetric crystals are at least two-fold degenerate, and their spin points in an arbitrary direction due to the freedom in describing the degenerate subspace. Computing spin-flip processes ab initio, especially in the presence of spin-orbit coupling (SOC) and spinor wave functions, remains an open challenge.

Here we present a first-principles method for computing the spin-flip e-ph coupling matrix elements and the $T_1$ spin-phonon relaxation times (SRTs). Our approach assumes no relationship between the matrix elements for spin-flip and momentum scattering, and treats spinor wave functions and SOC through fully-relativistic DFT and DFPT calculations [29]. These advances enable accurate calculations of SRTs and shed light on microscopic details of spin-phonon interactions. We apply our method to investigate SRTs in two key materials for spintronic and quantum technologies, silicon and diamond. Our predicted SRTs in silicon are in excellent agreement with experiment between 50–300 K, while in diamond, where SRT measurements are missing, we predict intrinsic-limit SRTs of roughly 0.5 ms at 77 K and 2 $\mu s$ at 300 K. In both materials, we find that spin-flip and momentum-scattering e-ph interactions differ widely and are not directly proportional, and the temperature dependence of the spin-flip and momentum relaxation times also differ greatly. Our work demonstrates a precise first-principles approach for computing SRTs, highlighting the limits of widely used simplified analyses and opening new avenues for microscopic understanding of spin dynamics.

Spin-flip interactions. In centrosymmetric materials, the Bloch states with band index $n$ and crystal momentum $\mathbf{k}$ can be decomposed into effective up and down spin states, denoted as $\uparrow$ and $\downarrow$, which diagonalize the $\hat{S}_z$ operator in the Kramers degenerate subspace [1, 2, 30]:

\[
\begin{align*}
\langle \mathbf{n}\mathbf{k}\uparrow | \hat{S}_z | \mathbf{n}\mathbf{k}\uparrow \rangle &= -\langle \mathbf{n}\mathbf{k}\downarrow | \hat{S}_z | \mathbf{n}\mathbf{k}\downarrow \rangle, \\
\langle \mathbf{n}\mathbf{k}\downarrow | \hat{S}_z | \mathbf{n}\mathbf{k}\uparrow \rangle &= 0.
\end{align*}
\]
The key ingredients for computing the SRTs are the spin-flip e-ph matrix elements [2],
\[ g^{\text{flip}}_{mn\nu}(k, q) = \langle mk + q | \Delta \hat{V}_{\nu q} | nk \rangle, \]
which quantify the probability amplitude to scatter from an initial Bloch state \( |nk\rangle \) to a final state \( |mk + q\rangle \) with opposite effective spin, by emitting or absorbing a phonon with mode index \( \nu \) and wave vector \( q \) due to the Kohn-Sham potential perturbation \( \Delta \hat{V}_{\nu q} \) [26], which is a \( 2 \times 2 \) matrix in spin space in the presence of SOC.

To compute the SRTs, we obtain the effective spin states and from them the spin-flip e-ph matrix elements \( g^{\text{flip}}_{mn\nu}(k, q) \) on fine BZ grids. We calculate the effective spin states from the spin matrix \( S(k) \), which provides a matrix representation of the spin operator \( \hat{g} e \times K \) and the Kohn-Sham potential perturbation \( \Delta \hat{V}_{\nu q} \), diagonal, with eigenvalues equal to the effective spin \( \Delta \hat{V}_{\nu q} \) [32]. The spin-flip e-ph matrix elements, \( g^{\text{flip}}_{mn\nu}(k, q) \), are then computed using Eq. (2) for all pairs of states with opposite effective spin.

**Interpolation.** Since DFPT calculations of \( \Delta \hat{V}_{\nu q} \) on the fine BZ grids needed to converge the SRTs are prohibitively expensive, we interpolate the spin-flip e-ph matrix elements \( S(k) \) and spin matrices using Wannier functions [33–35]. To obtain \( g^{\text{flip}}_{mn\nu}(k', q) \) at a desired pair of \( k' \) and \( q \) points in the BZ, we first apply the usual Wannier interpolation workflow [27, 36] to obtain the e-ph matrix elements \( g^{\text{flip}}_{mn\nu}(k', q') \) between states with arbitrary spins \( s \) and \( s' \). The e-ph matrix elements \( g^{\text{flip}}_{mn\nu}(k', q') \) coupling states with effective spins \( \sigma \) and \( \sigma' \) are then computed using the unitary matrix \( D_{k'} \) (the latter is obtained from Wannier interpolation of the spin matrix [31]):

\[ g^{\text{flip}}_{mn\nu}(k', q') = [D_{k' + q'}]_{s\sigma, ms} [g^{\text{flip}}_{mn\nu}(k', q')] [D_{k'}]_{ns', \sigma'}. \]

The spin-flip e-ph matrix elements are finally computed between all pairs of electronic states with opposite sign of the effective spin. Our interpolation scheme can accurately reproduce spin-flip e-ph matrix elements obtained by combining effective spin states with perturbation potentials computed directly with DFPT (see the Supplementary Material [37]), thus enabling precise calculations of SRTs.

**Spin relaxation times.** The band- and k-dependent spin-flip e-ph relaxation times, \( \tau^{\text{flip}}_{nk} \), are computed using lowest-order perturbation theory [2],
\[ \frac{1}{\tau^{\text{flip}}_{nk}} = \frac{4\pi}{\hbar} \sum_{\nu q} \left| g^{\text{flip}}_{mn\nu}(k, q) \right|^2 \left[ (N_{\nu q} + 1 - f_{mk + q}) \delta(\varepsilon_{nk} - \varepsilon_{mk + q} - \hbar \omega_{\nu q}) + (N_{\nu q} + f_{mk + q}) \delta(\varepsilon_{nk} - \varepsilon_{mk + q} + \hbar \omega_{\nu q}) \right], \]
where \( \varepsilon_{nk} \) and \( \hbar \omega_{\nu q} \) are the electron and phonon energies, respectively, and \( f_{mk} \) and \( N_{\nu q} \) the corresponding temperature-dependent occupations. Converging the BZ sum in Eq. (4) is challenging, especially since the spin-flip e-ph matrix elements \( g^{\text{flip}}_{mn\nu}(k, q) \) vary by several orders of magnitude throughout the BZ. We develop an importance sampling method to efficiently converge \( \tau^{\text{flip}}_{nk} \) (see the Supplemental Material [37]).

The temperature-dependent SRT, \( \tau_{\sigma}(T) \), is the main physical observable computed in this work. It is obtained as an ensemble average of the spin-flip relaxation times [2] by tetrahedron integration [38]:
\[ \tau_{\sigma}(T) = \left( \frac{1}{\tau^{\text{flip}}_{nk}} \right)^{-1} \left( \sum_{nk} \frac{1}{\tau^{\text{flip}}_{nk}} \left( \frac{d f_{nk}}{d E} \right) dk \right)^{-1} \left( \sum_{nk} \left( \frac{d f_{nk}}{d E} \right) dk \right), \]

**Numerical methods.** We apply our approach to investigate spin relaxation in silicon and diamond. We obtain their ground state and band structure using DFT with a plane-wave basis with the QUANTUM ESPRESSO code [42]. Briefly, we use relaxed lattice constants of 5.43 Å for silicon and 3.56 Å for diamond, together with a kinetic energy cutoff of 60 Ry for silicon and 120 Ry for diamond. We employ the PBEsol exchange-correlation functional [43] and fully-relativistic norm-conserving pseudopotentials [29] from Pseudo Dojo [44], which correctly include the SOC. We use DFPT [28] to compute the phonon dispersions and the perturbation potential, \( \Delta \hat{V}_{\nu q} \) in Eq. (2), on coarse q-point grids; our in-house developed PERTURBO code [45] is employed to compute the spin-dependent e-ph matrix elements on coarse BZ grids [46]. The DFPT calculations are done only in the irreducible q-point grid, following which we extend the coarse-grid e-ph matrix elements to the full q-point grid in PERTURBO by rotating the spinor wave functions with SU(2) matrices. The Wannier functions and spin matrices are computed with the WANNIER90 code [31] and employed in PERTURBO to interpolate the spin-flip e-ph matrix elements on fine BZ grids with up to 200 k-points to converge the SRTs. The spin quantization axis is chosen as the [001] direction.

**Temperature-dependent SRTs.** Figure 1(a) shows our calculated SRT as a function of temperature in silicon, which is in excellent agreement with experiments [39–41] at all temperatures between 50–300 K. For example, our calculated SRT at room-temperature is 4.9 ns, versus a 6.0 ns value measured by Lancaster et al. [41] [47]. The SRT in silicon exhibits an approximate \( T^{-3} \) temperature dependence; to explain its origin, we analyze in Fig. 1(a) the contributions from the three valley-dependent scattering processes, including the intravalley and so-called g and f intervalley processes, which correspond to scattering between valleys along the
The intravalley processes govern spin relaxation below 170 K, while the intervalley processes dominate above 170 K. We find SRTs of 540 \( \mu s \) at 77 K and 2.3 \( \mu s \) at 300 K; these values set an intrinsic limit due to phonons to the SRTs in diamond. The SRT exhibits a \( T^{-2} \) temperature dependence below \( \sim 170 \) K and a stronger \( T^{-5.5} \) trend above 170 K. This trend is in contrast with a previous prediction [20] of a \( T^{-5} \) temperature dependence throughout the entire temperature range and of an order-of-magnitude smaller SRT of 180 ns at room temperature. Ref. [20] assumed a direct proportionality between the spin-flip and momentum-scattering \( e\)-\( ph \) matrix elements, \( |g^{\text{flip}}(q)| \), with the spin-flip plus spin-conserving (i.e., momentum-scattering) \( e\)-\( ph \) matrix elements, \( |g^{\text{tot}}(q)| \), and resolves their ratio for different phonon modes. Depending on the phonon branch, we find that the spin-flip and momentum matrix elements can differ by several orders of magnitude, as we find for the longitudinal acoustic (LA) and longitudinal optical (LO) branches along \( \Gamma \rightarrow X \) and for the LO and for specific transverse optical (TO-1) and transverse acoustic (TA-2) branches along \( X \rightarrow K \rightarrow \Gamma \). For other phonon modes and BZ directions, the two quantities exhibit smaller — yet quantitatively important — differences. Only in specific cases the spin-flip and momentum-scattering interactions are nearly identical, as we find for the TO-2, TA-1 and LA branches along \( X \rightarrow K \rightarrow \Gamma \). These trends are common to silicon and diamond. Analogous results are found when analyzing various initial and final electronic states [37].

Lastly, we compare the spin-phonon and momentum relaxation times. The momentum relaxation time \( \tau_p \) is defined as the usual (spin-independent) \( e\)-\( ph \) relaxation time [26], thermally averaged using Eq. (5) to make the comparison meaningful. The conventional wisdom is that spin and momentum relaxation times are directly proportional [1, 22], an assumption that has been widely
used to analyze spin relaxation mechanisms in experimental data [49–54]. Figure 3 shows the temperature dependent spin and momentum relaxation times in silicon and diamond. In silicon, the SRT follows a $T^{-3}$ temperature dependence, whereas the momentum relaxation time follows a $T^{-2}$ trend. In diamond, the SRT makes a sharp transition from a $T^{-2}$ trend at low temperature to a stronger $T^{-5.5}$ trend above 170 K. In contrast, the momentum relaxation time exhibits a much weaker temperature dependence, roughly $T^{-1.5}$ at low temperature and $T^{-2.5}$ near room temperature.

There is no discernible direct proportionality between the spin and momentum relaxation times — rather, they both exhibit an approximate $T^{-n}$ temperature dependence, but with different values of the exponent $n$ (see Fig. 3). These differences originate from the different coupling strengths and phonon mode contributions, as we illustrate in Fig. 2. For example, we find that for momentum scattering in diamond the intravalley processes dominate over the entire temperature range up to 400 K, as opposed to just below 170 K as we show above for spin relaxation (see the Supplemental Material [37]). We conclude that a reliable analysis of SRTs needs atomistic calculations that take into account the different nature of the spin-phonon and momentum-scattering $e$-$ph$ interactions, using accurate spin-flip $e$-$ph$ matrix elements.

**Discussion.** Since SRT calculations involve a subtle interplay between spin-flip $e$-$ph$ matrix elements and phonons and electronic states, the relative magnitude of the spin-phonon interactions for different phonon modes is of paramount importance for accurate predictions. Our results show that the widely used proportionality between spin and momentum relaxation times can be inaccurate, highlighting the need for atomistic details such as the electronic wave function, spin texture, phonon modes and their mode-dependent spin-flip interactions. When these microscopic details are captured, as we have shown above, one can predict the SRTs within $\sim 10–20\%$ of experiment over a wide temperature range, and predict which phonon modes govern spin relaxation. While computing $e$-$ph$ interactions and carrier relaxation has become a main effort in first-principles calculations [55–59], SRT calculations are still in their infancy, and more work is needed to expand their scope beyond the EY mechanism discussed here.

**Conclusion.** In summary, we have developed a quantitatively accurate approach for computing spin-flip $e$-$ph$ interactions and SRTs due to the EY mechanism. The workflow proposed in this work is general — it can be adapted to different perturbation potentials, including perturbations from defects [60], through which one could study spin-flip and other defect-induced spin scattering processes. Our approach can be applied broadly to study spin relaxation in materials for spintronics and magnetism, and in topological materials. It can also be extended to treat spin states localized at ions or defects.
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diamond and \( 10 \times 10 \times 10 \) \( \mathbf{k} \)-point and \( \mathbf{q} \)-point grids in silicon.

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