Non-local correlation effects due to virtual spin-flip processes in itinerant electron ferromagnets

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We present an *ab initio* method for electronic structure calculations, which accounts for the interaction of electrons and magnons in ferromagnets. While it is based on a many body perturbation theory we approximate numerically complex quantities with quantities from time dependent density functional theory. This results in a simple and affordable algorithm which allows us to consider more complex materials than those usually studied in this context (3d ferromagnets) while still being able to account for the non-locality of the self-energy. Furthermore, our approach allows for a relatively simple way to incorporate self-consistency. Our results are in a good agreement with experimental and theoretical findings for iron and nickel. Especially the experimental exchange splitting of nickel is predicted accurately within our theory. Additionally, we study the halfmetallic ferromagnet NiMnSb concerning its non-quasiparticle states appearing in the bandgap due to spin-flip excitations.

I. INTRODUCTION

Even after the discovery of quantum mechanical exchange interactions as the reason for magnetic ordering, itinerant-electron magnetism including ferromagnetism of elemental Fe, Co, and Ni remained a challenging problem for decades [1–3]. The main issue is a coexistence of band (itinerant) and atomic (localized) features in the behavior of 3d-electrons in solids, something difficult to combine in a single theory [4]. The density functional theory (DFT), which was for a long time a method by default for a quantitative description of electronic properties of solids was applied to itinerant-electron magnets with great success [3] but atomic-like features such as remnants of atomic multiplet structure in some transition metal compounds were not easy to take into ac-The idea to combine the DFT with the dycount. namical mean-field theory (DMFT) [5-8] was an essential step to solve the problem. Ferromagnetic transition metals were one of the aims of this approach from the very beginning [7, 9, 10] and the results were very promising, especially in the description of their hightemperature magnetism and in the successful description of the famous 6 eV satellite in Ni [10]. Furthermore detailed calculations and comparison with the experimental data on angle-resolved photoemission [11–14] demonstrated that whereas the combination of the DFT and the DMFT seems to be sufficient for a quantitative description of Ni there are still remaining problems with Fe, despite an essential improvement of the description in the DFT+DMFT in comparison with the pure DFT approach. In particular, the DMFT being a local approximation predicts a wavevector-independent renormalization of effective masses with respect to the bare DFT values. Experimentally, it works quite accurate for Ni, a bit worse for Co and not accurate for Fe [14].

Another important group of itinerant-electron ferromagnets are weak itinerant ferromagnets such as ZrZn₂ [2] and half-metallic ferromagnets such as several Heusler alloys or CrO₂ [15]. The latter group of materials is especially interesting since for them many-body effects result not only in a renormalization and damping of electron quasiparticle spectrum but in appearance of a qualitative new feature, non-quasiparticle (or spin-polaron) states (NQPS) in a majority or minority electron energy gap [15–17]. These states were experimentally found in full Heusler alloy Co₂MnSi [18] and in CrO₂ [19]. The DFT+DMFT calculations can successfully describe NQPS at least at a qualitative level [15, 20]. At the same time, an appropriate quantitative description is possible only beyond the local approximation for the electron selfenergy, and in particular, it requires a correct description of the magnon dispersion [16, 17].

An important type of many-body effects in itinerantelectron magnets originates from the interaction of electrons with bosonic spin-flip excitations, both coherent (magnons) and incoherent (Stoner particle-hole) excitations [2]. While there has been a steady progress in understanding the properties of spin-flip excitations at a model level only little is known about microscopic details of their interactions with the electronic degrees of freedom in specific materials. The conventional DFT as well as the GW [21, 22] do not take these processes into account. The DFT+DMFT does take them into account

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but only in a local approximation whereas wavevectordependence of the corresponding contributions to the electron self-energy can be quite important, as we discussed above for the case of Fe. At the same time, the coupling has been directly experimentally detected [23–26] and is believed to crucially influence properties of magnetic materials leading to a number of remarkable phenomena. The latter include spin-dependent lifetime of excited electronic states [27] and inelastic electron transport [28]. Consequently, also the mean free path of excited electrons is spin dependent [29–31]. The electron-magnon interaction is also discussed as the moving force for the formation of Cooper pairs in certain high-temperature superconductors, in particular in the pnictide and cuprate families [32–35]. Furthermore, ultrafast magnetization switching phenomena in solids are believed to involve the coupling between the electronic and spin degrees of freedom as well [36, 37]. Last but not least, the coupling of magnons and electrons contributes to the renormalization of the electronic bands and influences the exchange splitting [38]. For the particular case of half-metallic ferromagnets, the effects of the electron-magnon interaction are systematically reviewed in Ref.[15].

Apart from the DFT+DMFT approach discussed above, the electron-magnon interaction can be treated within the many body perturbation theory (MBPT) [39, 40]. The MBPT relies on a perturbation expansion of the electronic self-energy based on the general Hedin equations [21]. MBPT calculations are computationally quite expensive, hence until recently only results from simple bulk systems are available [41, 42].

The approach presented in this work is based on the MBPT but avoids the numerical complexity by approximating some complex quantities with quantities from the time dependent DFT (TDDFT) [43-45], which are much easier to calculate. This allows to compute the influence of the electron-magnon interaction on the electronic structure for complex materials like half-metallic ferromagnets from first principles. This will allow us to consider NQPS taking into account non-locality of the spinflip contributions to the electron self-energy and thus going beyond the DFT+DMFT treatment of these states [15, 20]. Most recently, a method similar to ours including electron-magnon interactions based on MBPT was put forth [42]. However, while we opted to minimize the numerical burden of calculations (by using quantities from TDDFT and hence being able to account for more complex systems), they also included the GW term in their calculations leading to clearly improved results for the elemental 3d ferromagnets. We put forth another physically motivated simplification leading to a reduced computational cost of our calculations. In many cases, upon including the interaction, practically the entire spectral density of spin-flip processes is shifted from the Stoner energy window into the energy range of spinwaves [43]. This suggests that, unless the Landau damping itself is the subject of the investigation, it might be possible to replace the dynamical susceptibility with its "Heisenberg counterpart". We refer to this approach as "magnon-pole approximation". Its validity is discussed in detail and proven later in this paper.

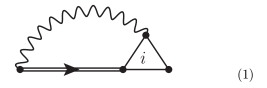
So far, our method is a one-shot method, meaning that we calculate the electronic structure of the materials in question and consequently perform the MBPT based algorithm to compute the self-energy representing the influence of the electron-magnon scattering. Van Schilfgaarde and coauthors [46] showed with a similar method, combining the DFT and the GW theory, that the results of this approach can depend on the starting electronic structure. It is shown in this work that one should ideally aim to use the results of the MBPT to construct a self-consistent cycle, making the final result independent of the starting Green function. Contrary to other methods based on the MBPT [41, 42], our approach offers a relatively simple way to incorporate a self-consistency. After the calculation of the electron-magnon self-energy, the renormalized Green function can be used to calculate new Heisenberg exchange parameters utilized in the magnon-pole approximation. Consequently, a new selfenergy can be calculated. The implementation of this numerical scheme is a current work in progress.

II. THEORY

A. Many body perturbation theory

Magnons are magnetic excitations, i.e. transitions to an excited state with a different value of the total spin projection. We will use this term for both coherent collective excitations (spin waves) and for incoherent spin-flip electron-hole pairs known as Stoner excitations [1–3]. Spin-waves are transverse fluctuations of magnetization that may be intuitively understood as a correlated precession of atomic magnetic moments. In the many body language they are formed due to the multiple scattering of particle and hole of opposite spin projection which are described, in a minimal set, as a summation of ladder diagrams [38, 47].

The interaction between magnons and electrons can be formulated in terms of an electron self-energy. Within the Hedin equations [21] the general expression for the self-energy reads



where the screened interaction W is drawn as wavy line, the many-body Green function G is represented by a double line and the three-leg vertex Γ (drawn as triangle)is

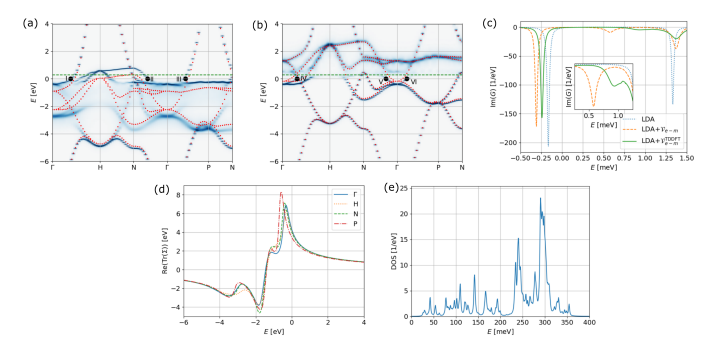
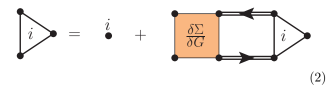


Figure 1: Effect of the electron-magnon interaction in bcc iron. (a) Electronic band structure of majority carriers under the influence of the electron-magnon interaction. The red dots represent the energy levels within the LDA while the blue background indicates the renormalized spectrum. The slope of the bands marked with black dots were compared to ARPES measurements in table I. The renormalized Fermi energy is represented by the dashed green line. (b) Same as (a) for minority carriers. (c) Comparison of the spectral functions obtained with $(LDA+\mathcal{V}_{e-m})$ and without $(LDA+\mathcal{V}_{e-m}^{TDDFT})$ the magnon-pole approximation at the Γ point for minority carriers. (d) Real part of the trace of the self-energy for different points in the Brillouin zone. (e) Magnonic density of states.

defined through the self-consistent equation



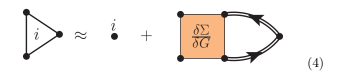
with the effective proper two-particle interaction (fourleg vertex) block given by the functional derivative

$$\frac{\delta \Sigma_{\alpha\beta}(1,2)}{\delta G_{\mu\nu}(4,3)}. (3)$$

In the latter expression the italic numbers represent the spatial and time variable 1 = (x, t).

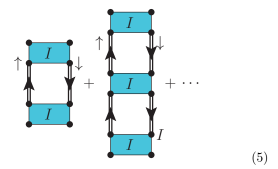
From now on we consider only ferromagnetic systems where charge fluctuations are decoupled from the transverse spin fluctuations. At the same time, they are coupled with longitudinal spin fluctuations [48, 49] but we will not consider these processes in the present paper. As long as the fundamental interaction (Coulomb potential in our case) is spin independent, even in the case of spin polarized systems, the screened interaction W cannot mediate a spin flip. Moreover, the screened interaction depends only on the charge part of the vertex function and therefore it is not affected by the electronmagnon interaction at all. Consequently, the only way

for the electron-magnon scattering to enter the theory is through the vertex correction in the self-energy relation 1. We use in our theory the three-leg vertex to first order

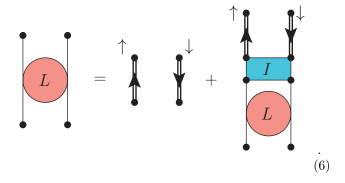


The first term in the latter equation, leading to the well known GW approximation [21, 22], does not involve electron-magnon scattering. Hence, for the remainder of this work, only the second term (linear in $\frac{\delta \Sigma}{\delta G}$) will be analyzed. As was shown in a number of preceding works [50, 51], $\frac{\delta \Sigma}{\delta G}$ can be divided into three classes of which only one type (diagrams reducible in the electron-hole channel) involves the scattering of an electron with a hole of opposite spin (i.e. involve magnons). Those diagrams

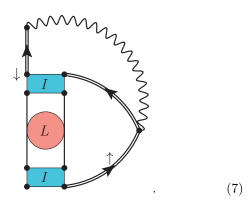
are of the form



where I are irreducible diagrams and the arrows represent the spin directions. The latter diagrams can be written compactly using the transverse two particle correlation function L defined through the Bethe-Salpeter equation



This allows to write the electron-magnon self-energy (often called GT contribution) as



The main drawback of the latter equation is its high numerical cost, especially for complex systems. Hence, more approximations are in order. First, I is assumed to be local

$$I_{\alpha\beta\mu\nu}(1,2,3,4) \approx I_{\alpha\alpha\mu\mu}(1,1,3,3)\delta(1-2)\delta(3-4).$$
 (8)

Similar approximations have been used successfully to calculate spin wave spectra [30, 38, 52]. Second, we also approximate the extra W in equation 7 with I. This

leads to a geometric series which can be summed. Furthermore, we make one more approximation in the spirit of Ng & Singwi [50] in starting the summation one term earlier. This allows to write the self-energy in terms of the susceptibility

$$\Sigma_{\alpha\beta}(1,2) \approx i\sigma_{\alpha\eta}^{i} I^{ij}(1,1,4,4) \chi^{jm}(4,6) I^{mn}(6,6,2,2) \sigma_{\eta'\beta}^{n} G_{\eta\eta'}(1,2)$$
(9)

as the susceptibility is given by the contraction of the two particle correlation function

$$\chi^{ij}(1,2) = -i\sigma^i_{\alpha\beta}\sigma^i_{\gamma\delta}L_{\alpha\beta\gamma\delta}(1,1^+,2^+,2). \tag{10}$$

In the latter equations we used a summation and integration convention: Each spatial or time variable (spin index) appearing only on one side of an equation is integrated (summed) over. The plus as superscript indicates that the time variable is modified by an infinitesimal ϵ : $1^+ = (\mathbf{x}, t + \epsilon)$.

B. Time dependent density functional theory

In order to arrive at a feasible computational scheme, we approximate the complex quantities from the many body perturbation theory by quantities from the time dependent density functional theory. The proper susceptibility can be obtained from the linear response TDDFT by means of the susceptibility Dyson equation [43, 48]

$$\chi_{\text{TDDFT}}^{ij}(\boldsymbol{x}_{1}, \boldsymbol{x}_{2}, E) = \chi_{\text{KS}}^{ij}(\boldsymbol{x}_{1}, \boldsymbol{x}_{2}, E) +$$

$$\sum_{m,n} \iint d^{3}x_{3} d^{3}x_{4} \chi_{\text{KS}}^{im}(\boldsymbol{x}_{1}, \boldsymbol{x}_{3}, E)$$

$$\left(K_{\text{xc}}^{mn}(\boldsymbol{x}_{1}, \boldsymbol{x}_{2}, E) + \frac{2\delta_{m0}\delta_{n0}}{|\boldsymbol{x}_{3} - \boldsymbol{x}_{4}|}\right) \chi_{\text{TDDFT}}^{nj}(\boldsymbol{x}_{4}, \boldsymbol{x}_{2}, E)$$
(11)

where the Kohn-Sham (KS) susceptibility is given by

$$\chi_{\text{KS}}^{ij}(\boldsymbol{x}_{1}, \boldsymbol{x}_{2}, E) = \sum_{mn} \sigma_{\alpha\beta}^{i} \sigma_{\gamma\delta}^{j} (f_{m} - f_{n})$$

$$\frac{\phi_{m\alpha}(\boldsymbol{x}_{1})^{*} \phi_{n\beta}(\boldsymbol{x}_{1}) \phi_{n\gamma}(\boldsymbol{x}_{2})^{*} \phi_{m\delta}(\boldsymbol{x}_{2})}{E - (E_{m\alpha} - E_{n\beta}) + i\epsilon}$$
(12)

with the Fermi-Dirac function f_n of the KS state $\phi_{n\alpha}$ at the energy $E_{n\alpha}$. In practice, the KS susceptibility and the exchange correlation kernel $K_{\rm xc}$ are computed interdependently from each other and, when combined in the susceptibility Dyson equation, yield slight deviation from the perfect Goldstone mode due to numerical inaccuracies. In our case, this deviation is of the order of few meV and can be removed (corrected) completely by imposing a suitable sum-rule on $K_{\rm xc}$. This issues were discussed carefully in [43] and [53].

To be fully consistent at this point, one also needs to

replace I with the exchange correlation kernel $K_{\rm xc}$ and replace the many body Green function with the KS Green function used for the calculation of the KS susceptibility (equation 12). This can be seen by comparison of equation 11 with the expression for the susceptibility derived in the framework of the MBPT 10 when I is assumed to be local. In this work we only use the exchange correlation kernel from the adiabatic local spin density approximation (ALSDA) [48]. This leads to the expressions

$$\begin{split} & \Sigma_{\uparrow}(1,2) = 2\mathrm{i} K_{\mathrm{xc}}^{+-}(1) \chi_{\mathrm{TDDFT}}^{-+}(1,2) K_{\mathrm{xc}}^{+-}(2) G_{\downarrow}(1,2) \\ & \Sigma_{\downarrow}(1,2) = 2\mathrm{i} K_{\mathrm{xc}}^{-+}(1) \chi_{\mathrm{TDDFT}}^{+-}(1,2) K_{\mathrm{xc}}^{-+}(2) G_{\uparrow}(1,2), \end{split}$$

$$(13)$$

which can also be written as

$$\Sigma_{\uparrow}(1,2) = iV_{e-m}^{-,\text{TDDFT}}(1^+,2)G_{\downarrow}(1,2)$$

$$\Sigma_{\downarrow}(1,2) = iV_{e-m}^{+,\text{TDDFT}}(1^+,2)G_{\uparrow}(1,2)$$
(14)

where

$$\mathcal{V}_{e-m}^{-,\text{TDDFT}}(1,2) = 2K_{\text{xc}}^{+-}(1)\chi_{\text{TDDFT}}^{-+}(1,2)K_{\text{xc}}^{+-}(2)
\mathcal{V}_{e-m}^{+\text{TDDFT}}(1,2) = 2K_{\text{xc}}^{-+}(1)\chi_{\text{TDDFT}}^{+-}(1,2)K_{\text{xc}}^{-+}(2)$$
(15)

represents a magnon mediated effective interaction. Analogous expressions have been derived in several reports using different theoretical approaches [50, 51, 54, 55]. Finally, the self energy is used to renormalize the electronic Green function obtained from the local density approximation (LDA) by means of the Dyson equation

$$G_{\alpha\beta}(1,2) = G_{\alpha\beta}^{0}(1,2) + G_{\alpha\mu}^{0}(1,3)\Sigma_{\mu\nu}(3,4)G_{\nu\beta}(4,2)$$
(16)

with the LDA Green function G^0 . The use of quantities readily available in the TDDFT substantially reduces the numerical burden, comparing to analogous expressions involving the screened Coulomb interaction [30, 38, 52, 56, 57] and allows to address large systems like NiMnSb and in the future two dimensional heterostructures.

In practice, the calculation of the electron-magnon interaction with the TDDFT susceptibility is still problematic as the susceptibility is needed for an immense number of energy points. To reduce the number of points the fluctuation-dissipation theorem can be utilized [47]. After the calculation the spectral weights

$$C_{\text{TDDFT}}^{\pm}(\boldsymbol{r}, \boldsymbol{r}', \boldsymbol{k}, E) = i\theta(E) \left[\chi_{\text{TDDFT}}^{\pm}(\boldsymbol{r}, \boldsymbol{r}', \boldsymbol{k}, E + i\epsilon) - \chi_{\text{TDDFT}}^{\pm}(\boldsymbol{r}', \boldsymbol{r}, \boldsymbol{k}, E + i\epsilon)^{*} \right]$$
(17)

along a path parallel to the real axis, the TDDFT susceptibility at arbitrary energies in the complex plane can be calculated by simple integration of the spectral representation of the susceptibility

$$\chi_{\text{TDDFT}}^{\pm}(\boldsymbol{r}, \boldsymbol{r}', \boldsymbol{k}, E) = \int \frac{dE'}{2\pi} \frac{\mathcal{C}^{\pm}(\boldsymbol{r}, \boldsymbol{r}', \boldsymbol{k}, E')}{E - E' + i\epsilon} - \int \frac{dE'}{2\pi} \frac{\mathcal{C}^{\mp}(\boldsymbol{r}', \boldsymbol{r}, \boldsymbol{k}, E')}{E + E' - i\epsilon}$$
(18)

However, due to the high numerical cost of the TDDFT susceptibility, this approach is only viable for fixed \boldsymbol{k} and energies close to the Fermi energy. Hence, the magnon-pole approximation (discussed in the chapter II C) was used for all following results unless otherwise noted.

With the magnon propagator V_{e-m} playing the role of the effective interaction, equations 14 convey a compelling physical picture of the electron-magnon scattering in solids. Electrons (holes) in a given state can emit a magnon and decay into another electron (hole) state of opposite spin providing energy and momentum differences between the initial and the final particle states are accounted for by this magnon. Furthermore, due to the prevalence of up-to-down magnon processes over the down-to-up ones, in strong ferromagnets, effectively, only down electrons and up holes interact with the spin-flip excitations, respectively loosing and gaining energy in this process. The above observations allow us to justify the deployment of the TDDFT quantities in the MBPT scheme.

A magnon corresponds to the transverse fluctuations of the magnetization δm . In the spirit of the TDDFT, these fluctuations give rise to the fluctuating exchangecorrelation potential (the transverse magnetic field in the case of a collinear ground state and the ALSDA) while the potential is given by the exchange-correlation kernel K_{xc} , such that $\delta B_{xc}^{\pm} = K_{xc} \delta m^{\pm}$. We note that the very same relationship governs calculations of the enhanced susceptibility in the susceptibility Dyson equation, eq. 11. Thus, the abstract concept of electronmagnon scattering is reinterpreted as the interaction of the electron with the space- and time-dependent fluctuating exchange-correlation field associated with the magnon [58]. In the case of the TDDFT this field is straightforwardly given as $K_{\rm xc}\delta m^{\pm}$ where $K_{\rm xc}$ naturally plays the role of the interaction vertex in our equations. Recall that equivalent interpretation appears in quantum electrodynamics [59] where the interaction of an electron and photon can be understood as the interaction of the electron with the electromagnetic field corresponding to this photon.

Let us, however, recall that the equations 20 are obtained upon including only the minimal set of diagrams involving scattering of electron-hole pairs of opposite spins. Therefore, while the exchange of virtual magnons appears to be a dominating contribution of spin fluctuations on the band structure in itinerant magnets, it is necessarily only an approximate, albeit compelling model.

C. Magnon-pole approximation

While the Kohn-Sham susceptibility (equation 12) consists of a broad spectrum of spin flip excitations located at the energy scale of the exchange splitting, the TDDFT susceptibility (equation 11) develops new low energy sin-

gularities corresponding to spin waves, i.e. coherent precession of the atomic magnetic moments. Interestingly, in most cases, practically the entire spectral density is shifted from the Stoner energy window into the energy range of the spin-waves. This suggests that, unless the Landau damping itself is the subject of the investigation, it might be possible to replace the dynamical susceptibility by its "Heisenberg counterpart" χ^H with singularities located at the undamped spin-wave energies and spatial dependence corresponding to the rigid tilt of the atomic magnetic moments μ :

$$\left[\chi_{\rm H}^{-1}\right]_{ij} = (2g\mu_i)^{-1} \left(z\delta_{ij} + g\mu_j^{-1}J_{ij} - g\delta_{ij}\mu_j^{-1} \sum_{\ell} J_{i\ell} \right)$$
(19)

Here, the exchange interactions J were calculated from the first principles magnetic force theorem [60] and the Lande-factor $g \approx 2$ was used. The relation between the full expression for the dynamical susceptibility and its Heisenberg-like form (19) is discussed in detail in Ref. [48]. We show that in many cases the use of this Heisenberg-like susceptibility instead of the TDDFT susceptibility in the expression for the self-energy changes the results only marginally. At the same time, as computations of the TDDFT susceptibility are computationally demanding, this "magnon-pole-approximation" results in a relatively inexpensive numerical scheme. We also note that the magnon-pole approximation is equivalent to the assumption of a strong ferromagnet. Using this approximation, the self-energy is given by

$$\Sigma_{\uparrow}(1,2) = i\mathcal{V}_{e-m}^{-}(1^{+},2)G_{\downarrow}(1,2)$$

$$\Sigma_{\downarrow}(1,2) = i\mathcal{V}_{e-m}^{+}(1^{+},2)G_{\uparrow}(1,2)$$
 (20)

where

$$\begin{split} \mathcal{V}_{e-m}^{-}(1,2) &= 2K_{\text{xc}}^{+-}(1)\chi_{\text{H}}^{-+}(1,2)K_{\text{xc}}^{+-}(2) \\ \mathcal{V}_{e-m}^{+}(1,2) &= 2K_{\text{xc}}^{-+}(1)\chi_{\text{H}}^{+-}(1,2)K_{\text{xc}}^{-+}(2). \end{split} \tag{21}$$

D. Goldstone mode, double counting and self-consistency

We conclude this section with a short comment on double counting which is a typical problem in approaches combining DFT with more sophisticated methods like the dynamic mean field theory or the MBPT [12, 41]. The problem arises on two different levels in methods presented in the literature. First, the introduction of two different MBPT terms can lead to double counting as discussed in [41]. The authors include (part of) the GW term [53] for their susceptibility to restore the correct Goldstone mode and then study the impact of the electron-magnon interaction with the same diagrams we use in this work. This leads to double counting if the diagram of second order in I is included in the calculation of the electron-magnon scattering

self-energy (see the text above equation 13). In fact, the second order term alone is unphysical as it already includes double counting [41]. To strictly avoid double counting, one has to start from the third order in I by subtracting the second order term from the geometrical series of ladder diagrams. However, it was shown in [41, 42] that this leads to the violation of the causality manifesting itself in the wrong sign of the imaginary part of the self-energy. This, in turn, can be corrected by the inclusion of numerically expensive GW term itself [42], which, however, does not contribute to the spin-dependent electron scattering and is not considered in this work. For the materials studied in this work, the electron-magnon interaction exerts essentially the same effect on the electronic structure as in [41] and DMFT methods [7, 11–14]. Hence, the double counting issue does not seem to be decisive here. Furthermore, in our case the exclusion of the second order term would also lead to a significant increase in the numerical cost of our scheme (especially in combination with the magnon-pole approximation discussed below). Hence, we choose to include the second order diagram. As discussed in section II.B, the resulting full geometric series of ladder diagrams corresponds to a compelling physical picture of an electron interacting with the fluctuating Kohn-Sham potential associated with the magnon. Concerning the fact that your scheme is not self-consistent and we take the Kohn-Sham band structure as the starting point, we find the approximation physically acceptable.

Second, double counting may arise due to the fact that it is unknown which many body effects are already included in the exchange correlation potential of the DFT. While the double counting of the first kind can be taken into account exactly (at least in principle), this is not the case for the second kind. In the literature these double counting issues are often assumed to be small due to the fact that the DFT potentials are based upon the homogeneous electron gas which includes only little correlation effects at the relevant densities [41]. Other publications utilize a heuristic double counting correction [12]. In this work we resort to the first option and assume the double counting corrections are small enough to neglect them.

Finally, we note that our method is a one-shot scheme, i.e. it is not self-consistent. This can cause similar methods to lead to different results due to slightly different LDA Green functions as was also recognized when the LDA Green function was renormalized with the GW term [46]. The obvious way to overcome this problem is to impose some kind of self consistency, which is a work in progress. In our method, a self-consistent scheme can be constructed by using the calculated self-energy to calculate new interaction parameters J which are then used to calculate the renormalized Heisenberg susceptibility, leading to a new self-energy. The implementation of this scheme is still a work in progress.

III. RESULTS AND DISCUSSION

A. bcc iron

The spectrum of iron under the influence of the electron-magnon interaction is presented in figure 1 a and b. Generally, the electron-magnon interaction leads to a broadening of the electron states as well as a renormalization of the electron energies and the appearance of additional peaks. Our result agrees qualitatively with recently published results for iron using an the MBPT method [41, 42] and the DMFT methods [7, 11–14]. All of the aforementioned works report larger broadening for majority spin carriers, especially around E = -2 eV across the whole spectrum. In the minority spin channel we observe an additional peak at Γ for $E \approx 0.5$ eV, which does not appear in any of the other works. The spectral weight of this peak is however very small and hence, it might be due to a slightly different LDA Green function compared to [41]. It is certainly not caused by the magnon-pole approximation as will be shown in the following. Regarding the renormalization of the electronic energies compared to the LDA solution, our results show the same trend that was found in [41] and in the GT results of [42]. The different absolute values are partly due to the magnon-pole approximation and can also arise due to a slightly different LDA Green functions.

While the majority spin electrons are shifted to higher energies, the minority bands remain mostly at the same energies. Due to the fact that several majority bands cross the LDA Fermi energy, the renormalizatation is expected to also have an impact on the Fermi energy. The change of the Fermi energy can be calculated from the renormalized density of states (DOS) by "filling up" the states until the correct number of electrons is reached. It needs to be mentioned at this point that the calculation of the renormalized DOS is only feasible when the self-energy is assumed to be constant in k, which is not quite the case for iron as will be discussed in more detail later. Hence, the resulting changes of the Fermi energy (and the magnetic moment) for the case of iron should be viewed as an approximation. The green dashed line in figure 1 a and b indicates the new Fermi level which lies about 0.27eV above the LDA Fermi energy. The magnetic moment per atom is reduced to $\mu = 2.05 \mu_{\rm B}$ giving a slight deviation from the experimental result of $\mu \approx 2.2 \mu_{\rm B}$ [61] which is accurately predicted within the LDA. The renormalization of the Fermi energy and the magnetic moments is significantly higher for iron than for any other material studied here. It may be speculated that a self-consistent procedure is more important in this case.

Note that the spectrum of spin up electrons is mostly affected below the Fermi energy while for spin down electrons the opposite trend can be observed. This is caused by the imaginary part of the self-energy being approximately zero above (below) the Fermi energy for minority (majority) spin carriers. Physically, this asymmetry arises from the extreme case of a strong ferromagnet at

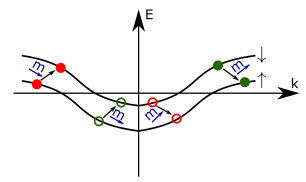


Figure 2: Different quasiparticle transitions involving magnons in a simplified band structure. A collinear magnet without spin-orbit interaction at low temperatures is considered. As there are no thermally excited magnons, the magnon emission processes dominate. In the strong ferromagnets, the magnons are mainly associated with the decrease of the angular momentum (they are, effectively, up-to-down spin-flips). From left to right: A majority electron absorbs a magnon (blue arrow labeled m) and gets excited into an empty minority spin state. A majority hole is excited into a minority hole by emitting a magnon. A minority hole decays into a majority hole by absorbing a magnon. A minority electron decays into a majority state and emits a magnon in the process. At low temperatures, the processes colored green are dominant while the processes colored red are negligible. There are further thinkable spin-dependent processes: the emission and absorption of spin excitations which increase the total spin angular momentum (involving, effectively, down-to-up spin-flips). At low temperatures, they could be emitted by an up electron or down hole. In the ferromagnets discussed in this paper, these so called anti-Stoner processes feature negligible spectral weight and are not considered.

zero temperature. In figure 2 all possible quasiparticle transitions in a strong ferromagnet are schematically drawn. Above the Fermi energy a minority spin electron can decay into a majority spin state by emitting a magnon. The opposite process of a majority spin electron being excited into a minority state does not occur as it would need a (e.g. thermally excited) magnon to account for spin conservation. The corresponding argumentation applies for holes below the Fermi energy.

The damping of spin-waves through hybridization with the Stoner continuum is an important mechanism in materials with itinerant electrons. Especially in iron this effect was shown to lead to "spin-wave disappearance" [43]. Thus, although most of the spectral weight of the magnetic spectrum lies in the spin-wave region, the assumption of sharp magnon peaks is generally not fulfilled throughout the Brillouin zone. Hence, we study the influence of the magnon-pole approximation for a small energy range at the Γ point and minority carri-

ers, as shown in figure 1 c. The LDA result is compared with the renormalized Green functions computed within the magnon-pole approximation (LDA+ \mathcal{V}_{e-m}) and with the TDDFT susceptibility (LDA+ $\mathcal{V}_{e-m}^{\text{TDDFT}}$). The previously discussed peak is visible for both Green functions renormalized with the TDDFT susceptibility and its Heisenberg counterpart. The position of the peaks differs slightly and the peaks calculated with the TDDFT susceptibility are slightly broader. However, we conclude that the major effects of the renormalization are properly included within the magnon-pole approximation.

Over recent years several experimental studies of the band structure of iron were conducted [13, 14, 62]. In all these studies the effective mass renormalization $\frac{m_{\alpha}^{*}}{m_{0\alpha}^{*}}$ was measured by means of angle resolved photoemission spectroscopy (ARPES). Using DFT and ARPES quasiparticle energies, Sanchez-Barriga et. al. [13, 14] derive an energy dependence of the self-energy from which they calculate the effective mass renormalization and compare to DMFT results. They find that in bcc iron the effective mass renormalization depends strongly on the position in the Brillouin zone. However, the DMFT cannot account for a k dependence in the self energy and hence gives the same result independent of the position in the BZ. While they show that this is a good approximation for Ni the situation is different in the case of Fe. Our results for the trace of the self-energy calculated for iron (figure 1) d) and nickel (figure 3 e) at different positions in the BZ show that for nickel the self-energy is practically constant while for iron a clear dependence on the position in the BZ can be observed. This is also manifested in the values for the effective mass renormalization which we find to vary between 1.54 at the N point and 2.25 at the Γ point for the Σ_4 band. Hence, our results confirm that the non-local character of electron correlations in bcc iron is essential for the proper band structure description. The physical origin of this difference between Fe and Ni will be discussed below, at the end of the next subsection. Schaefer et. al. [62] compare the slopes of bands at the Fermi energy instead (i.e. for constant energy instead of constant k). As reference they use their own DFT calculations which, although qualitatively similar, differ from our results. To eliminate the dependence on the DFT reference we compare the velocity of the renormalized quasiparticle obtained in [62] directly from the experimental data, cf. table I. We calculate the velocity by taking the derivative of the quasiparticle energy (i.e. the position of the peak maximum) with respect to k. Apart from point I, the electron-magnon interaction reduces the quasiparticle velocity compared to the LDA which leads to a better agreement with the experimental data.

B. fcc Ni

Nickel is an example of a material not well described by the LDA [10]. Here, the most prominent features are the missing satellite peak at 6 eV binding energy, a far too

	Experiment [62] (eVÅ)	Our result (eVÅ)	
		$\left ext{LDA} + \mathcal{V}_{e-m} \right $	LDA
Г-Н (І)	1.12	2.2	2.2
Spin $\uparrow \Gamma$ -N (II)	1.16	1.2	3.5
Γ -P (III)	1.4	2.3	4.3

Table I: Velocity of quasiparticles $v = \frac{\partial \varepsilon}{\partial k}$ at the Fermi energy in bcc iron from the experimental study [62] compared with our results. The labels next to the direction refer to the points in the spectrum (figure 1 a and b).

	LDA	$\mathrm{LDA} + \mathcal{V}_{e-m}$	GWT	LDA+DMFT	exp.
L_3	0.71	0.31	0.37	~0.4	$0.31 {\pm} 0.03$
X_2	0.7	0.23	0.31		~ 0.2

Table II: Comparison of exchange splittings in nickel (in eV) with the GWT approach [42], the LDA+DMFT [14] and experimental results [63, 64].

large exchange splitting and a too broad valence band. These discrepancies are believed to be a result of many-body effects, which cannot be described within the LDA, in particular those arising from the coupling to spin fluctuations. The DFT+DMFT solves the problem and allows to reach a very good agreement with the experiment eliminating all three discrepancies [10, 11, 14]. At the same time, the DMFT takes into account different kinds the many-body processes. It is instructive to separate the effects of the electron-magnon interaction, which we will do here, without local approximation for the electron self-energy.

We find that the 6 eV satellite of Ni is not originated by coupling to spin-flip excitations. Also, the narrowing of the valence band width is not a consequence of the electron-magnon scattering. In fact, the bandwidth is slightly increased for the renormalized Green function, cf. figure 3 a and d. Thus, from these three problems of the LDA, only one, namely, the overestimate of the spin splitting, is related to electron-magnon processes and is fully grasped in our approach.

Generally, the majority spin bands are strongly damped by the electron-magnon interaction as can be seen by the blue background in figure 3 compared to the LDA solution indicated by red filled dots. The minority spin carriers in contrast are hardly influenced by the electron-magnon interaction, cf. figure 3 b. This is due to the spin asymmetry of the density of states, as also reported in [41]. We note that the exchange splitting is strongly reduced by a new peak in the majority spin channel appearing throughout the whole spectrum close to the Fermi energy visible in figures 3 a and 3 d. Our results for the exchange splitting at the L and Xpoints are in excellent agreement with ARPES results, as shown in table II. The shift of majority electron states towards the Fermi energy as a consequence of the electron-magnon interaction is reflected clearly in the

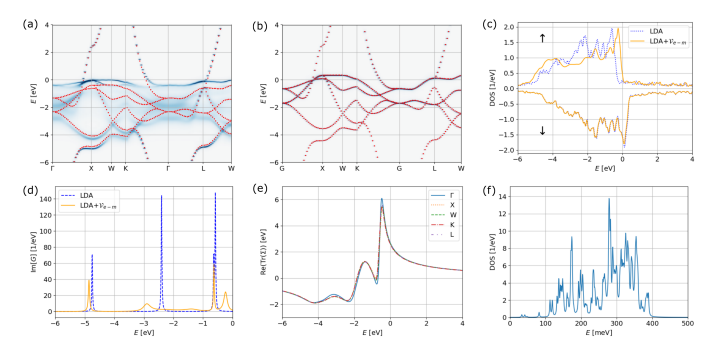


Figure 3: Effect of the electron-magnon interaction in fcc nickel. (a) Electronic band structure of majority carriers under the influence of the electron-magnon interaction. The red dots represent the energy levels within the LDA while the blue background indicates the renormalized spectrum. (b) Same as (a) for minority carriers. (c) Electronic density of states within the LDA and the LDA under the influence of the electron magnon scattering (LDA+ \mathcal{V}_{e-m}). (d) Spectral function at the L point from the LDA and the renormalized Green function. (e) Real part of the trace of the self-energy for different points in the Brillouin zone. (f) Magnonic density of states.

DOS, as shown in figure 3 c. However, in the ARPES experiments no additional peak (compared to LDA results) was found. The reason for this can be seen in figure 3 d which shows the spectral weight (which is proportional to the imaginary part of the Green function) at the L point for the LDA and renormalized Green functions. The LDA results include a peak at $-2.5 \,\text{eV}$ which is damped by the strong electron-magnon interaction to such an extent that it can hardly be identified as a well defined quasiparticle any more. Consequently, our results only feature three clear peaks in agreement with the ARPES studies.

Similar to iron, a shift of the majority spin bands towards higher energies can be observed in figure 3 a. However, here the renormalized bands are still mostly located below the Fermi energy. Consequently, the renormalization has practically no effect on the Fermi energy (it shifts by less than 30meV) and the magnetic moment (it decreases by approximately $0.072\mu_{\text{B}}$).

As was already mentioned at the end of the previous section, the self-energy in nickel is much less dependent on the wavevector than in iron, in agreement with the previous conclusions based on the comparison of DFT+DMFT data with the experiment [10, 14]. The difference might be associated to an essential difference of exchange interactions and thus magnon spectra between bcc Fe and fcc Ni. Already from the values of

the spin-wave stiffness constant it is clear that magnons in Fe are much softer than in Ni despite the fact that the magnetic moments of the atoms are almost four time larger in the former case than in the latter. This is related with two circumstances. First, Friedel oscillations of exchange parameters are much more pronounced for Fe than for Ni [65, 66]. The reason for this is that Ni is almost half-metallic, with almost fully occupied majorityspin band whereas the Friedel oscillations require metallicity for both spin projections. Second, the magnetic interactions in Fe are frustrated, with the ferromagnetic e_q electron subsystem and the antiferromagnetic t_{2q} - t_{2q} contributions to the exchange parameters [67]. On the opposite, magnetism of Ni is mostly determined by the nearest-neighbor ferromagnetic interactions without any essential competing interaction.

Let us conclude this section with a short comment on the electron-magnon scattering in the spin-polarized uniform electron gas at densities corresponding to the 3d transition metals as it can provide us with an additional insight into the emergence of the magnetic order in these systems. A simple model of these magnets involves two parabolic bands split by a uniform exchange-correlation magnetic field B_{xc} . The Fermi energy $E_{\rm F}$ and B_{xc} are chosen such that the charge density equals to the one of the valence band of a 3d system and the magnetic moment assumes the corresponding value. This model yields a qualitatively correct picture of the magnons in itinerant

systems [2] with a magnon band featuring the Goldstone mode and a quadratic dispersion for small wave vectors, and interacting with the Stoner continuum. Following our scheme, also the self-energy arising due to the exchange of virtual magnons can be computed. It turns out, however, that its value is much larger than $E_{\rm F}$ and would destroy the magnetic order thus yielding a qualitatively incorrect picture of magnetism in this model. The failure is attributed to the unrealistically large exchange splitting B_{xc} , and correspondingly the exchange-correlation kernel K_{xc} , necessary to induce the magnetic moment according to the Stoner criterion $(K_{xc}\chi_0 = 1)$. In real 3d transition metals, the emergence of the magnetic order arises due to the large value of the non-interacting susceptibility χ_0 being in turn proportional to the large density of states in the partially localized 3d band. Thus, the real K_{xc} coupling is small and the self-energy, as expected, turns out to be a correction to the band structure.

C. NiMnSb

NiMnSb is seen as a prototype example for a half-metallic ferromagnet (HMF), i.e. it is conducting for one spin direction and a semiconductor for the other [15, 68]. In the case of NiMnSb the LDA density of states near the Fermi energy is zero for the minority spin channel as depicted in figure 4 a. Prediction of the half-metallicity in this compound [68] opened the field of half-metallic ferromagnetism.

The use of the magnon-pole approximation is particularly justified in half-metallic ferromagnets. The gap in the electronic spectrum at the Fermi level causes the Stoner excitations to emerge at energies much larger than typical for spin-waves. The latter are therefore Stoner undamped and contain the majority of the spectral weight [15, 69].

The emergence of non-quasiparticle states (NQPS) in the spin gap is a trademark of half-metallic ferromagnets. The physical picture of such states is that spin down states directly above the Fermi energy can be seen as a superposition of spin up states and virtual magnons [15, 17, 20]. Contrary to quasiparticle states, which arise from the poles of the Green function, NQPS are caused by its branch cuts [17]. There is no spectral weight of such states directly at the Fermi energy but it increases drastically at energy scales of characteristic magnon frequencies. The spectrum of minority spin carriers, cf. figure 4 b, shows a strong broadening of electronic states above the Fermi energy. Across the whole spectrum a dispersionless peak appears approximately 150 meV-300 meV above the Fermi energy. The magnonic density of states [70, 71], cf. figure 4 c confirms a high number of magnon states within that energy region (the electronic DOS of majority carriers is approximately constant in the relevant energy window, cf figure 4 a). Note that this density of states looks much simpler than that for Fe (Fig. 1 e) and Ni (Fig. 3 f), with just a few well-pronounced main peaks. Similar to the case of nickel, the self-energy of NiMnSb depends only weakly on the position in the BZ which makes our results quite close to results obtained from the DFT+DMFT [20]. This is a general feature of the self-energy for most materials as the magnon energies are small compared to the typical electronic bandwidth. As we already discussed, bcc Fe seems to be a counterexample. However, in this case its ferromagnetism is related to \mathbf{e}_g states only, and one can assume that instead of the total bandwidth it is the width of this peak which matters [67, 72].

Hence, the electronic density of states may be calculated under the assumption that the self-energy is independent of \boldsymbol{k} hence lowering the numerical burden of the calculation. Our results, presented in figure 4 a, show a broadening of the sharp peaks in the DOS caused by the broadening of the electron states through the electron-magnon interaction. In addition, a peak in the DOS emerges in the gap in the minority spin channel due to the NQPS formation.

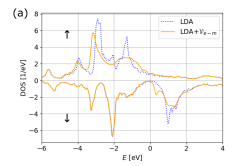
Similar to the situation in nickel, the Fermi energy as well as the magnetic moments of the system remain practically unchanged through the renormalization (the Fermi energy increases by 86meV and the total magnetic moment increases by $0.001\mu_{\rm B}$).

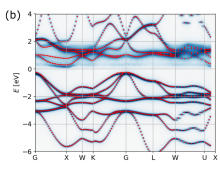
IV. SUMMARY AND OUTLOOK

In this work we presented an ab initio method to account for the influence of non-local correlations due to spin-flip processes on the electronic structure of ferromagnets. It is based on the MBPT in the formalism of Hedin [21] but avoids its main disadvantage, i.e. the high numerical complexity, as the main quantities used are approximated using TDDFT. This methods gives very similar results to other more complex methods [41, 42] for bcc iron and fcc nickel. Most notably, the predicted exchange splitting in fcc nickel is closer to the experimental value than those of other comparable methods. The lower numerical cost allows to study complex materials as the half-metallic ferromagnet NiMnSb. In the latter material, the electron-magnon interaction causes the appearance of non-quasiparticle states in the bandgap. Contrary to DMFT, we are able to account for the nonlocality of the self-energy. Extensions of the theory towards antiferromagnets and 2D systems as well as the influence of paramagnons are a current work in progress.

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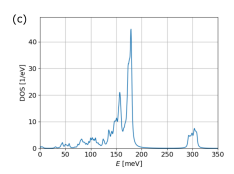


Figure 4: Effect of the electron-magnon interaction in NiMnSb. (a) Electronic density of states within the LDA and the LDA under the impact of the electron magnon scattering (LDA+ \mathcal{V}_{e-m}). (b) Electronic band structure of minority carriers under the influence of the electron-magnon interaction. The red dots represent the energy levels within the LDA while the blue background indicates the renormalized spectrum. (c) Magnonic density of states.

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