Spin-fluctuation and spin-relaxation effects of single adatoms from first principles

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Abstract

Single adatoms offer an exceptional playground for studying magnetism and its associated dynamics at the atomic scale. Here we review recent results on single adatoms deposited on metallic substrates, based on time-dependent density functional theory. First we analyze quantum zero-point spin-fluctuations (ZPSF) as calculated from the fluctuation-dissipation theorem, and show how they affect the magnetic stability by modifying the magnetic anisotropy energy. We also assess the impact of ZPSF in the limit of small hybridization to the substrate characteristic of semi-insulating substrates, connecting to recent experimental investigations where magnetic stability of a single adatom was achieved for the first time. Secondly, we inspect further the dynamics of single adatoms by considering the longitudinal and transverse spin-relaxation processes, whose time-scales are analyzed and related to the underlying electronic structure of both the adatom and the substrate. Thirdly, we analyze spin-fluctuation modes of paramagnetic adatoms, i.e., adatoms where the Stoner criterion for magnetism is almost fulfilled. Interestingly, such modes can develop well-defined peaks in the meV range, their main characteristics being determined by two fundamental electronic properties, namely the Stoner parameter and the density of states at the Fermi level. Furthermore, simulated inelastic scanning tunneling spectroscopy curves reveal that these spin-fluctuation modes can be triggered by tunneling electrons, opening up potential applications also for paramagnetic adatoms. Lastly, an overview of the outstanding issues and future directions is given.

Single adatoms deposited on surfaces have become a prominent playground where theory and experiment can explore hand-in-hand a large variety of physical phenomena ranging from spin-excitations [1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11, 12, 13, 14] to magnetic exchange interactions [2, 15, 16, 17, 18, 19, 20], quantum spin decoherence [21, 22, 23], interplay between magnetism and superconductivity [24, 25, 26, 27, 28, 29, 30], or the Kondo effect [31, 32, 33, 34, 35, 36, 37], among many others. Virtually all these effects arise from the intricate interplay between the degrees of freedom of the adatom — charge, spin or orbital angular momentum — and the electron and phonon bath of the substrate, a subject of heavy and ongoing investigation. Noteworthily,
magnetism plays a central role in fueling the interest for these tiny objects, given that they represent the ultimate limit in the context of bit miniaturization in data storage devices. Therefore, understanding the above mentioned phenomena is of capital importance both for fundamental physics and for potential technological applications. In this context, the field has witnessed an enormous push forward after the advent of spin-polarized scanning tunnelling microscopy (STM) and inelastic electron tunnelling spectroscopy (IETS). These experimental techniques, occasionally used in combination with X-ray magnetic circular dichroism (XMCD) \cite{38, 39} and electron paramagnetic resonance (EPR) \cite{22}, allow to monitor the dynamical regime by, e.g., measuring atomic spin-excitations \cite{1, 3, 40, 8} and quasiparticle interferences \cite{17}, accessing spin-relaxation and decoherence times \cite{41, 22, 19} and even resolving highly dynamical processes like the reading and writing of magnetic information \cite{42}.

From the theoretical point of view, early simulations based on density functional theory (DFT) \cite{43, 44, 45} served for boosting and creating a huge enthusiasm in the field by predicting gigantic local magnetic moments of diverse transition-metal (TM) adatoms, including 4d and 5d elements that are nominally nonmagnetic in bulk. In addition, the understanding and interpretation of features present in IETS experiments, such as the characteristic steps in the conductance spectrum \cite{46}, greatly benefited from studies based on quantum spin models, e.g., by showing the importance of inelastic excitation channels involved in the process (see, e.g., Refs. \cite{47, 6, 48, 49, 50, 51}). In the last few years, the development of a real-space spin-dynamics technique within time-dependent DFT (TDDFT) \cite{7, 52} has proven to be a powerful tool for characterizing the central dynamical quantity, namely the spin-excitation spectrum, which contains information about the exchange of energy and angular momentum with the substrate electrons that form the environment of the magnetic adatoms \cite{7, 52}. Importantly, a quantitative analysis of the role played by the spin-orbit coupling (SOC) in determining the resonance frequency of spin-excitations and their reaction to magnetic fields has been successfully given within this approach \cite{53}. Going one step beyond, the use of many-body perturbation theory (MBPT) in connection with TDDFT has also made possible to establish a direct quantitative comparison to the $dI/dV$ (differential tunneling conductance) curves measured in IETS experiments \cite{54}.

One of the most fundamental requirements for a successful usage of single adatoms for technological purposes is the stability of its magnetic moment. This, in turn, depends crucially on the so-called magnetic anisotropy energy (MAE), an energy barrier generated by the SOC that protects and stabilizes the direction of the local magnetic moment against possible fluctuations of the spin, e.g., of thermal origin. Noteworthily, XMCD and IETS assert that several 3d transition metal (TM) adatoms can possess a substantial MAE of few meV: examples include Fe and Co on Pt(111) \cite{55, 9}, Fe on Cu(111) \cite{8} as well as on Cu$_2$Ni/Cu(111) \cite{56} and CuNi \cite{3}, and Co on MgO(100) \cite{57}, among others. Strikingly, however, these large MAEs do not directly translate into an stable magnetic moment, given that the above mentioned systems behave as paramagnetic entities when measured by means of STM (see, e.g., Refs. \cite{58, 59, 16}), thus implying the existence of a mechanism that destroys the magnetic stability locally. Going one step further, the case of 4d and 5d adatoms is even more striking, given that they have so far exhibited no clear magnetic signal even when subjected to the static magnetic field of an XMCD experiment, in notorious disagreement with theoretical predictions \cite{60}. Up to date, the only single adatom that has experimentally exhibited magnetic remanence is Ho on MgO/Ag(100) \cite{38, 42}, i.e., a
4f rare-earth element with a very large local magnetic moment ($\sim 10 \, \mu_B$) deposited on a thin insulating layer.

In view of this rich scenario, a central question arises: what is the mechanism leading to the apparent instability of the magnetic moment of an adatom? Given that temperature effects are not the principal cause of the magnetic instability [58], one must turn to other destabilizing effects. Recent efforts to address this far-reaching issue have been mainly based on symmetry arguments applied to model Heisenberg Hamiltonians, whereby the switching probability of the magnetic moment is calculated by means of a master equation (see e.g. Refs. [61, 62, 63, 38]). These arguments apply to a well-localized magnetic moment of either integer or half-integer spin, which is justified in the weak coupling regime to the substrate. If the coupling is strong, as it is generally the case for metallic substrates, the orbitals responsible for the magnetic moment of an adatom hybridize with the surface conduction electrons, giving rise to an itinerant electron picture with fractional local magnetic moments (see, e.g., Refs. [8, 45, 7, 52, 53]). In the latter case, the proper description of the itinerant character is needed for addressing the subject of magnetic stability.

The present review focuses on the aforementioned strongly-hybridized metallic systems. In this context, zero-point spin-fluctuations (ZPSF) become a key dynamical aspect which are intimately connected to spin-excitations through the so-called fluctuation-dissipation theorem [64]. ZPSF are quantum fluctuations present even at absolute zero temperature that can crucially affect the magnetic properties of itinerant electron magnets. The importance of spin-fluctuations was first realized in the early studies of weak bulk ferromagnets [65, 66, 67, 68] and has subsequently been verified in the context of DFT and TDDFT too [69, 70]. In this review, we consider the concept of ZPSF in single adatoms and quantify their impact on the magnetic stability of the series of 3d and 4d TM adatoms deposited on metallic substrates. We show that ZPSF are of the order of the local magnetic moment itself, an extremely large value that has deep effects on the MAE, which can be reduced by more than 50% with respect to the static value calculated by standard DFT. We further characterize the three fundamental ingredients that determine the magnitude of the ZPSF, namely the (i) local magnetic moment, (ii) SOC and (iii) electron-hole Stoner excitations. Based on our findings, we develop a simple diagram where the ZPSF of an arbitrary adatom are classified according to these ingredients, offering practical guidelines for achieving magnetic properties which are robust against fluctuations.

Apart from protecting the magnetic stability, in order to achieve the ultimate goal of a technologically useful single adatom it is mandatory to master the dynamical properties associated to its spin. For example, fast spin-dynamics can be useful when the goal is to transfer magnetic information from or to the adatom, while slow spin-dynamics are desirable if the aim is to store magnetic information. What are the typical time scales involved, and how does the environment affect them? Interestingly, the lifetime of an atomic spin-excitation can be experimentally accessed from the width of the step observed in IETS $dI/dV$ measurements, being typically of the order of picoseconds [8, 57, 3, 71]. Throughout this review, we quantitatively connect this lifetime to electronic properties of adatoms and their hybridization with the substrate, showing that the single-particle spin-flip excitation channel is the main driving mechanism behind it. In tandem, recent experimental developments yielded another way to measure spin-relaxation times using an all-electrical pump-probe technique [41], which has been so far applied to adatoms.
deposited on insulating substrates [57, 22, 72]. Based on the ab initio perspective, here we show that, in order to successfully perform such experiments in metallic systems, a measuring scheme based on ultrafast techniques able to record relaxation times of the order of femtoseconds would be needed.

Although magnetic adatoms have by far attracted most of the interest in this field, nominally non-magnetic adatoms can also exhibit interesting fingerprints of magnetism and thus be potentially useful for technological purposes. This is the last main topic addressed in the present review, where we study what are the requirements for observing well-defined features in the spin-excitation spectrum of this type of adatoms, i.e., paramagnetic spin-excitations (PSE). These are the analogous of so-called paramagnons first proposed by Doniach in 1967 [73] and first measured in bulk Pd nearly 50 years later by Doubble and co-workers [74] (see also Ref. [75] for recent calculations). Upon reducing the dimensionality of the system, here we show that PSE can be strongly enhanced due to the modified interplay between the two fundamental electronic properties involved, namely the Stoner exchange interaction and the adatom’s density of states (DOS) at the Fermi level. In addition, the ab initio analysis reveals that PSE are highly sensitive to externally applied magnetic fields and, furthermore, can exhibit a singular enhancement when the field approaches a critical regime. Finally, we assess the impact of PSE on the dI/dV signal as measured in IETS experiments, identifying clear signatures of magnetic response that allow to distinguish these type of excitations from, e.g., phonons.

The review is organized as follows. In Sec. 1, we sketch the theoretical formalism used to calculate the quantities presented throughout the review. In Sec. 2 we begin by presenting the ground state properties of a set of 3d and 4d TM adatoms that will be the main focus of the review. In Sec. 3 the concept of ZPSF is presented as well as an analysis of its magnitude and its effect on inducing magnetic instability via the renormalization of the MAE barrier. In Section 4, we focus on discussing atomic time scales for the relaxation time of the spin. Finally, in Sec. 5 we illustrate the concept of paramagnetic spin-fluctuation modes and possible means of probing it experimentally. Conclusions and outlook are given in Sec. 6, while appendices A and B contain derivations on the Bloch and Landau-Lifshitz-Gilbert models, respectively.

1 Theoretical approach

In this section we settle the theoretical background used throughout the review. For this, let us consider the linear response of a ferromagnetic system to an externally applied time-dependent perturbation,

\[ \delta \mathbf{m}(r; t) = \int dr' \int dt' \chi(r, r'; t - t') \delta \mathbf{V}(r'; t'). \]

Above, \( \delta \mathbf{m} = (\delta m_x, \delta m_y, \delta m_z, \delta n) \) and \( \delta \mathbf{V} = (\delta B_x, \delta B_y, \delta B_z, \delta V) \), with \( \delta m_i \) and \( \delta B_i \) respectively the components of the vector spin density and external magnetic field, while \( \delta n \) and \( \delta V \) are the charge density and external scalar potential, respectively. Working in frequency space and defining atomic-like quantities by integrating out the spatial dependence over atomic sites [52], the above expression takes the simplified form

\[ \delta \mathbf{m}(\omega) = \chi(\omega) \delta \mathbf{V}(\omega). \]
For a single magnetic atom, the quantity $\chi$ in the above equations is a 4×4 tensor that couples in general all components of the spin and charge responses with each other. If SOC is weak, however, the full response decouples into a longitudinal and transverse part [76]. This approximation is justified for the systems investigated in this review since the off-diagonal sectors of the susceptibility tensor are small in comparison to the diagonal ones. Then, assuming that the perturbation is purely of magnetic origin (i.e. $\delta V = 0$), the change of the spin magnetic moment length is described by

$$\delta m_z(\omega) = \chi_\parallel(\omega) \delta B_z(\omega). \quad (3)$$

Above, $\chi_\parallel(\omega)$ denotes the longitudinal spin-susceptibility. This quantity is determined by excitations between electrons with same spin state, given that it involves the Pauli matrix $\sigma_z$ that is diagonal in spin basis [77]. On the other hand, the change of the transverse spin components can be compactly described using the circular combinations $m_\pm = m_x \pm i m_y$ and $B_\pm = B_x \pm i B_y$, yielding for the + component

$$\delta m_+(\omega) = \chi_+(\omega) \delta B_+(\omega). \quad (4)$$

Above, $\chi_\pm(\omega)$ denotes the transverse spin-susceptibility which, contrary to $\chi_\parallel(\omega)$, is determined by transitions that flip the spin state of the electrons due to the transverse Pauli spin matrices involved, which are off-diagonal in spin space [7, 52, 53]. We note that the full transverse response is given by

$$\chi_\perp(\omega) \equiv 2(\chi_+(\omega) + \chi_-(\omega)). \quad (5)$$

The spin-susceptibility is closely connected to the ZPSF via the so-called fluctuation-dissipation theorem [64]. This theorem relates the variance of the spin-fluctuations along a given direction, $\xi^2_i$, to the imaginary part of the corresponding spin-susceptibility, $\text{Im} \chi_i(\omega)$,

$$\xi^2_i = -\frac{1}{\pi} \int_0^\infty d\omega \text{Im} \chi_i(\omega), \quad (6)$$

with $i = \parallel, \perp$.

The calculation of the ZPSF requires therefore a realistic and accurate description of the spin-susceptibility of the adatom. This can be achieved by employing the time-dependent generalization of DFT, namely TDDFT [78, 79]. The general expression for the adatom’s enhanced spin-susceptibility within the TDDFT framework is given by [7, 52, 53, 80, 54, 81]

$$\chi_i(\omega) = \chi_{i\text{KS}}^\parallel(\omega) \left(1 - \frac{U_i}{\chi_{i\text{KS}}^\parallel(\omega)} \right), \quad (7)$$

with $i = \parallel, \pm, \mp$. Above, $\chi_{i\text{KS}}^\parallel(\omega)$ is the bare KS spin-susceptibility, i.e. the standard response function in linear response theory. For the analysis provided in this review, we calculate the different components of $\chi_{i\text{KS}}^\parallel(\omega)$ using the Korringa-Kohn-Rostoker Green function (KKRGF) method employing the atomic sphere approximation with full charge density [82]. For details on the formalism, we refer the reader to Refs. [52, 53].

Finally, we note that $U_i$ in Eq. 7 denotes the interaction kernel, which in this work is treated in the adiabatic local spin-density approximation [83]. In the longitudinal case, the kernel
Figure 1: Calculated ground state electronic properties of single adatoms. (a) Spin magnetic moments for several 3\textit{d} transition metal adatoms deposited on Ag(100) and Cu(111) denoted by circles (blue) and triangles (red), respectively. (b) Same as (a) but for 4\textit{d} adatoms. (c) Atom-projected total electronic DOS for 3\textit{d} adatoms deposited on Ag(100). The majority and minority spin channels are denoted by solid (positive) and dashed (negative) lines, respectively. The vertical line denotes the Fermi level.

\( U_{||} \) contains two contributions, namely the Coulomb term as well as the exchange-correlation (XC) term responsible for the many-body effects \([70, 84]\). In the transversal case, the Coulomb term drops out leaving only the XC term. Importantly, the transverse kernel \( U_{\pm} = U_{\mp} \) can be connected to the magnetic moment \( m_0 \) and the XC magnetic field \( B_{xc} \) via sum-rules that allow an accurate description of the dynamical regime: for details, we refer the reader to Refs. \([7, 52, 53]\).

2 Preliminaries: ground state properties of 3\textit{d} and 4\textit{d} magnetic TM single adatoms deposited on metallic substrates

The main focus of this review is a set of 3\textit{d} and 4\textit{d} adatoms deposited on the metallic Ag(100) and Cu(111) substrates. Before tackling their dynamical properties, we first analyze two ground state quantities, namely the spin magnetic moment, denoted as \( m_0 \), and the electronic DOS. Let us begin with Fig. 1(a) and 1(b), where the calculated \( m_0 \) is depicted. This figure shows that all the considered adatoms develop large magnetic moments of more than 2\( \mu_B \). Furthermore, \( m_0 \) acquires non-integer values, indicating the itinerant character of the adatom’s \( d \)-electrons induced by the hybridization with the electrons of the metallic substrate. This feature is confirmed by the DOS, which is displayed in Fig. 1(c) for the specific case of 3\textit{d} adatoms deposited on Ag(100). This figure shows that the \( d \)-state peaks, so-called virtual bound states, are substantially broadened (between \( \sim 0.1 \) eV and \( \sim 1 \) eV depending on the adatom), a well-known consequence of hybridization with the substrate \([44, 8, 53]\). Another property evidenced by Figs. 1(a,b) is that the first of the atomic Hund’s rules is closely fulfilled, i.e. the half filled \( d \)-shell elements develop largest magnetic moments, case of Cr and Mn for 3\textit{d}, Mo and Tc for 4\textit{d}. Finally, Figs. 1(a,b) show that the choice of metallic substrate and surface orientation does
not substantially affect the spin magnetic moment of the adatom, implying that the substrate’s symmetry plays only a minor role in this context. We note that these ground state properties are consistent with the original works by Dederichs and co-workers [43, 44, 45], as well as with more recent studies [7, 52, 53, 80, 54, 81].

3 Zero-point spin-fluctuations

In this section we analyze the ZPSF (Eq. 6) associated to the magnetic single adatoms presented in Sec. 2. Two fundamentally different contributions can be distinguished: the longitudinal ($\xi^2_\parallel$) and transverse ($\xi^2_\perp$) components of the ZPSF. In Table 3 we have listed the calculated magnitude of both quantities. Importantly, these values show that the longitudinal contribution is approximately an order of magnitude smaller than the transverse one: the longitudinal component reaches a maximum of $\sqrt{\xi^2_\parallel} = 0.38 \mu_B$ for Mo on Ag(100), while $\sqrt{\xi^2_\perp} > 1 \mu_B$ for virtually all the adatoms. This feature reflects the fact that it is much easier to alter the direction of the magnetic moment than its size. Thus, given that the major contribution to the fluctuations comes from the transverse component, in the following we will focus on its analysis and its effects on the magnetic stability.

3.1 Connection to spin-excitation spectra

The integrand of Eq. 6 defining the ZPSF is composed by the imaginary part of the enhanced spin-susceptibility, a quantity that displays so-called spin-excitations associated to the damped precessional motion of the spin-moment [7, 52, 53]. Calculated transverse spin-excitation and spin-fluctuation spectra for TM adatoms deposited on Ag(100) are shown in Fig. 2. Let us first focus on Figs. 2(a,c), which display $\text{Im} \chi_\perp(\omega)$ for 3$d$ and 4$d$ adatoms, respectively. These figures reveal the existence of a large peak in the meV range for all adatoms, corresponding to a spin-excitation. Note that the associated resonance frequency is dependent on the MAE, which is in turn settled by the SOC [53]. The largest resonance frequency of $\sim 6$ meV pertains to the Fe adatom, while the rest of adatoms display the peak at $\lesssim 4$ meV; in the particular cases of Ti, V, Mn, Cr and Mo, we find the spin-excitation at extremely small frequencies, $\lesssim 0.5$ meV, implying that the net effect of SOC is very weak in these adatoms. Figs. 2(a,c) display yet another feature, namely the width of the spin-excitation peak, which is linked to the amount of electron-hole Stoner excitations near the Fermi level [80]: Ru and, to some extent also Nb and Tc, display large widths as compared to the rest of adatoms, in particular those that peak

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<td>$\sqrt{\xi^2_\parallel \text{[Ag(100)]}}$</td>
<td>0.09</td>
<td>0.11</td>
<td>0.06</td>
<td>0.07</td>
<td>0.14</td>
<td>0.18</td>
<td>0.21</td>
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<tr>
<td>$\sqrt{\xi^2_\parallel \text{[Cu(111)]}}$</td>
<td>0.12</td>
<td>0.14</td>
<td>0.12</td>
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<td>$\sqrt{\xi^2_\perp \text{[Ag(100)]}}$</td>
<td>2.05</td>
<td>2.50</td>
<td>3.21</td>
<td>3.25</td>
<td>3.16</td>
<td>1.63</td>
<td>0.92</td>
<td>2.41</td>
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<td>1.88</td>
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<td>$\sqrt{\xi^2_\perp \text{[Cu(111)]}}$</td>
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<td>2.80</td>
<td>2.31</td>
<td>2.56</td>
<td>3.02</td>
<td>2.57</td>
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Table 1: Calculated longitudinal and transverse contribution to the ZPSF (units of $\mu_B$).
Figure 2: Calculated spin-excitation spectra and ZPSF. (a) and (c) Density of transverse spin-excitations as given by $\text{Im} \chi_{\perp}(\omega)$ for selected 3$d$ and 3$d$ adatoms on Ag(100), respectively. Insets address different frequency regions where various resonance frequencies are located. (b) and (d) Calculated magnitude of the mean value of transverse ZPSF for the adatoms considered in (a) and (c), respectively. Solid lines depict the evolution of the value as a function of the upper boundary of the integral of Eq. 6, while the horizontal dashed lines represent the converged value.

below 1 meV.

We next turn to the magnitude of the transverse ZPSF (Eq. 6), as displayed in Figs. 2(b,d) for 3$d$ and 4$d$ adatoms on Ag(100), respectively; the solid lines depict the evolution of the mean value, $\sqrt{\xi^2_{\perp}}$, as a function of the upper boundary of the frequency integral, whereas the converged value is marked by the horizontal dashed lines. As already pointed in the discussion of Table 3, $\sqrt{\xi^2_{\perp}}$ is of the order of the Bohr magneton; the elements with largest values are Cr, Mn, Fe and Tc, with $\sqrt{\xi^2_{\perp}} \sim 3\mu_B$, whereas in Zr and Co this value is reduced by more than 50%. As the calculations demonstrate, the main contribution to the integral of Eq. 6 comes from the spin-excitation peak in the meV region, which represents between 70% and $\sim100\%$ of the total depending on the adatom.

Figs. 3(a) and 3(b) show an instructive comparison between the magnitude of the transverse ZPSF and the local magnetic moments of the adatoms. These figures demonstrate that $\sqrt{\xi^2_{\perp}}$ (blue stars) is always an appreciable fraction of $m_0$ (blue circles); one finds $\sqrt{\xi^2_{\perp}} \sim m_0/2$ for Mn and Cr, while $\sqrt{\xi^2_{\perp}} \sim m_0$ for Ti, Fe, Zr, Nb and Ru, revealing that the latter suffer from strong deviations of the direction of the local magnetic moment. We also note that both $m_0$ and $\sqrt{\xi^2_{\perp}}$ follow the evolution predicted by Hund’s rules, i.e. adatoms with nearly half-filled $d$-shells have largest values: Cr and Mn among 3$d$, and Mo and Tc among 4$d$. This trend is clearly fulfilled in the case of the magnetic moment as already mentioned in Sec. 2, while the evolution of the fluctuations shows some exceptions like Fe and Tc. Noteworthily, a similar relationship between magnetic moment and fluctuations has been very recently found also for bulk transition metals [85]. In addition, Fig. 3 also reveals that the spin-fluctuation-to-magnetization ratio (SFMR) is generally larger in 3$d$ adatoms than in 4$d$. As a final remark, let us note that Mo, which is the only 4$d$ adatom deposited on Ag(100) that shows an experimentally measurable magnetic signal [86], has by far the lowest SFMR among 4$d$ elements.

Figs. 3(a) and 3(b) also display an extension of the analysis to the same set of adatoms deposited on Cu(111), which exhibit essentially the same features as on Ag(100). This again indicates that symmetry plays a minor role in these largely hybridized metallic systems with strong itinerant
character. The only noteworthy difference is that the SFMR of Cr and Mn is somewhat lower than what is expected from the trend of Fig. 3; we will discuss the origin of this feature shortly. Let us also note that Co adatoms show a very large SFMR in both Ag(100) and Cu(111) (see Fig. 3). Incidentally, Co is known to behave as a Kondo system when deposited on various metallic substrates [87, 88, 89]. Since both ZPSF and the Kondo effect are driven by spin-flip excitations with substrate electrons (see discussion below and Fig. 4), it is tempting to associate both effects. However, only a more advanced theory capable of describing such many-body effects can prove this hypothesis.

### 3.2 Interpretation in terms of the Landau-Lifshitz-Gilbert model

In the following, we turn to identify the fundamental factors that determine the magnitude of the transverse ZPSF. For this purpose, we consider the Landau-Lifshitz-Gilbert (LLG) equation [90]. The LLG is widely employed for characterizing the spin-dynamics of macroscopic magnetic systems, and more recently also for microscopic systems [80, 53] as it allows to accurately reproduce the \textit{ab initio} calculations by extracting the relevant parameters (see also Ref. [91] for a derivation of the quantum counterpart of the LLG equation). The details of the model are worked out in Appendix B. Here, let us recall that within the LLG model, the imaginary part of the transverse spin-susceptibility takes the form of an skewed Lorentzian, i.e.

\[
\text{Im}\chi_{\lambda\mu}^{\text{LLG}}(\omega) = \frac{M\gamma}{2(1 + \eta^2)} \cdot \frac{\eta\omega}{(\omega - \omega_0)^2 + (\eta\omega_0)^2}.
\]  

(8)

The parameters entering this model are the Gilbert damping, \(\eta\), which determines the width of the spin-excitation peak and is dominated by Stoner excitations, and the resonance frequency, \(\omega_{\text{res}} = \omega_0\sqrt{1 + \eta^2} = \gamma B_{\text{eff}}/\sqrt{1 + \eta^2}\), where \(\gamma\) is the gyromagnetic ratio and \(B_{\text{eff}}\) an effective magnetic field whose magnitude is determined by the strength of SOC (and of the external field, if considered); a detailed discussion can be found in, e.g., Ref. [53].

The integral of Eq. 6 can be calculated analytically within the LLG model and yields immediately the expression for the transverse ZPSF:

\[
\xi^2_{\lambda\mu}^{\text{LLG}} = \frac{M\gamma}{\pi(1 + \eta^2)} \left(\frac{\eta}{2} \log \left(\frac{x^2 + \eta^{-2} + 1}{(\eta^{-2} + 1)^2} \right) + \vartheta(x, \eta)\right).
\]  

(9)
The background shows a 2D plot of the SFMR as a function of the damping and the resonance frequency in the LLG model (Eq. 9). Circles (black) and squares (red) denote various adatoms deposited on Ag(100) and Cu(111), respectively, where the parameters $\eta$ and $\omega_0$ (see discussion of Eq. 8) have been extracted from a fit to the \textit{ab initio} spin-susceptibility.

Above, $\psi(x, \eta) = \arctan(x - \eta^{-1}) - \arctan(x + \eta^{-1}) + 2 \arctan \eta^{-1}$ and $x = \omega_c/\eta\omega_0$, with $\omega_c$ a cutoff frequency to be converged.

Eq. 9 provides an interpretation for the magnitude of the fluctuations in terms of the physical parameters of the LLG model, which are in turn related to the electronic structure of the adatoms. In particular, three major ingredients come into play: $m_0$, $\omega_0$ and $\eta$. First of all, Eq. 9 reveals that $\sqrt{\xi^2_{\text{LLG}}}$ is proportional to $\sqrt{m_0}$, which already accounts for the rough proportionality displayed by the \textit{ab initio} calculations (see Fig. 3). Moreover, this relationship further indicates that fluctuations are relatively weaker for adatoms with large magnetic moments, namely the nearly half-filled $d$-shell elements. Interestingly, the simple relationship of Eq. 9 allows to analyze the SFMR, $\sqrt{\xi^2_{\text{LLG}}}/\gamma m_0$, as a function of $\eta$ and $\omega_0$ by setting the standard value for the gyromagnetic ratio $\gamma = 2$; the map obtained by doing so is shown in the background of Fig. 4. This figure evidences that the SFMR is very much enhanced by the damping since $\sqrt{\xi^2_{\text{LLG}}}/\gamma m_0$ varies by almost 70% in the range of values considered for $\eta$. In contrast, Fig. 4 shows that the transverse ZPSF are reduced for large resonance frequencies, but the induced variation is much less important than that of the damping. In order to quantitatively connect these features to \textit{ab initio} calculations, we have extracted the parameters $\eta$ and $\omega_0$ by fitting $\text{Im}\chi^{\text{LLG}}(\omega)$ to the calculated spin-susceptibility (e.g., Figs. 2(a,c)) for all adatoms. In this way, we locate the position of each adatom on the map of Fig. 4, as depicted by the circles and squares for the case of the Cu(111) and Ag(100) substrates, respectively. The resulting distribution evidences that the origin of the large SFMR found in the case of Ti, Co, Ru and Nb on both substrates, as well as Fe and Zr on Ag(100), is mainly due to the large damping factors of these adatoms, $\eta \gtrsim 0.5$, while this tendency is only slightly modified by the position of the resonance frequency.
We note that a remarkable feature is revealed when $\eta$ tends to zero, i.e.,

$$\lim_{\eta \to 0} \sqrt{\xi_{\text{LLG}}^2 / \gamma m_0} = 1 \mu_B^{1/2} \Rightarrow \lim_{\eta \to 0} \sqrt{\xi_{\text{LLG}}^2} = \sqrt{\gamma m_0}. \quad (10)$$

The expression above shows that the transverse ZPSF remain finite even in the extreme case of a spin-excitation with vanishing width. This intrinsic contribution is required by Heisenberg’s uncertainty principle, since different spin-operators do not commute with each other [92], thus setting a lower boundary for the fluctuations. This is indeed the case found for Mn, Cr and Mo adatoms, which, despite their small resonance frequency, their tiny damping ($\eta \lesssim 0.05$) makes $\sqrt{\xi_{\text{LLG}}^2 / \gamma m_0}$ approach the intrinsic minimum value for these elements. Interestingly, we note that Eq. 10 represents a sensible limit to give a rough estimate for the ZPSF of adatoms deposited on insulating and semiconducting substrates, since the substrate hybridization giving rise to the damping tends to be far smaller on those type of substrates as compared to the metallic ones considered in this review. Let us illustrate this notion by considering the important example of a Ho atom deposited on ultrathin MgO(100) layers grown on Ag(100), which experimentally exhibits a stable magnetic moment for time-periods larger than $10^3$ s [38]. From Hund’s rules the Ho adatom should have a magnetic moment of $10\mu_B$, as found experimentally [42]. Under this assumption, a quick estimate using Eq. 10 reveals that the magnitude of the ZPSF in Ho should be $\sim 4.5\mu_B$. This, in turn, means that the resulting SFMR is nearly a factor of 2 smaller than in the elements with lowest SFMR studied in this review (see Eq. 13 and Fig. 5), implying that the relative impact of the quantum spin-fluctuations for Ho on MgO/Ag(100) is particularly weak.

### 3.3 Impact on the magnetic anisotropy energy

As a last step in the analysis of this section devoted to spin-fluctuations, we turn now to analyze the effects of the transverse ZPSF on the magnetic stability of the adatoms. For this purpose, we estimate how the fluctuations affect the MAE, the quantity defining the height of the energy barrier when the magnetic moment is made to rotate away from its easy axis. We note that the MAE is closely connected to the resonance frequency of the spin-excitations (see, e.g., Ref. [53]) that give rise to the primary contribution to the transverse ZPSF; most importantly, the energy scale of both quantities is the same (meV).

We consider the following expression for the MAE for uniaxial systems,

$$E_a(\theta) = K \frac{(m \cdot \hat{e}_z)^2}{m^2} = K \cos^2 \theta, \quad (11)$$

where $K$ is the so-called anisotropy constant. As a consequence, the energy barrier between the magnetic moment pointing along the $z$ axis ($\theta = 0$) and a perpendicular axis ($\theta = \pi/2$) is given by $E_a(\theta = 0) - E_a(\theta = \pi/2) = K$, as it is schematically illustrated in Fig. 5(a). In the spirit of the spin-fluctuation theory of Moriya [65], we now let magnetic moment fluctuate around its equilibrium value, i.e. $m^2 \rightarrow (m_0 \hat{e}_z + \sum_\perp \xi_\perp)^2$. Introducing this term into the definition of $E_a(\theta)$, we obtain a renormalized expression for the MAE, i.e.

$$E_a(\theta, \xi_\perp^2) = \frac{K (m_0^2 \cos^2 \theta + \xi_\perp^2 \sin^2 \theta)}{m_0^2 + 2\xi_\perp^2}. \quad (12)$$
Figure 5: (a) Schematic illustration of the renormalization of the MAE induced by the transverse ZPSF. K and K' respectively represent the static and renormalized anisotropy constant (see Eq. 13). (b) and (c) Calculated anisotropy constants for 3d and 4d adatoms, respectively. Blue hexagons and leftward triangles denote K and K' for adatoms on Ag(100), respectively, while red squares and upward triangles denote K and K' for adatoms on Cu(111), respectively.

In practice, this implies that the energy barrier gets effectively reduced by the transverse ZPSF,

$$E_a(\theta = 0, \xi_{\perp}^2) - E_a(\theta = \pi/2, \xi_{\perp}^2) = K \left(1 - \frac{3\xi_{\perp}^2}{m_0^2 + 2\xi_{\perp}^2}\right) \equiv K'. \quad (13)$$

The above is characterized by a modified anisotropy constant K', also illustrated in Fig. 5(a).

In Figs. 5(b) and 5(c) we show the calculated values for the anisotropy constant, K, as well as for the reduced one, K': the former has been evaluated by band energy differences following the magnetic force theorem [93], while the latter has been calculated using the values of the magnetic moments and transverse ZPSF of the adatoms. These figures show a large and generalized reduction of the MAE, which goes beyond estimates that do not take into account fluctuation effects [53]. In cases where the ZPSF are larger than $m_0$ (mostly 4d adatoms), the magnetic moment direction becomes destabilized, as reflected by the sign change in the renormalized MAE.

Let us pay special attention to the well-studied case of Fe on Cu(111), where the MAE is reduced from $K = -4.73$ meV to $K' = -0.32$ meV. Interestingly, this means that ZPSF considerably remedy the disagreement with the experimental MAE, which is nearly $\sim -1$ meV [8, 94]. More importantly, this example shows that the renormalization of the MAE due to the ZPSF has the correct order of magnitude. Focusing next on the 4d elements Ru and Nb on Ag(100), we find that the reduction of the MAE predicted by the calculations is notably large, i.e. $\gtrsim 90\%$. It is noteworthy that these two cases have been thoroughly studied experimentally, which have concluded that there is no sizable magnetic signal in the systems [60, 95]. This suggests that the ZPSF might also play a major role in the experimentally observed absence of stable magnetism.
4 Spin-relaxation times

While the analysis of Sec. 3 deals with the magnitude of the fluctuations, this section instead focuses on the time scales. For this purpose, we consider two particular relaxation processes of magnetic single adatoms, namely the longitudinal and transverse spin-relaxations characterized by the relaxation times $T_{\parallel}$ and $T_{\perp}$, respectively. Physically, $T_{\parallel}$ characterizes the relaxation of the size of the adatom’s spin magnetic moment while $T_{\perp}$ describes its damped precessional motion. In the following, we analyze each of this processes separately.

4.1 Longitudinal component

In essence, the longitudinal spin-susceptibility $\chi_{\parallel}(\omega)$ (see Eq. 7) describes the ability of the system to continuously modify the size of its magnetic moment by an externally applied time-dependent magnetic perturbation along the magnetization direction. The dynamics of this process can be phenomenologically studied in terms of the longitudinal Bloch equation, which yields the following form for the enhanced spin-susceptibility [96] (see Appendix A),

$$\chi^{\text{Bl}}(\omega) = \frac{\chi_0^{\text{Bl}}}{1 - i\omega T_{\parallel}}.$$  (14)

Above, $\chi_0^{\text{Bl}}$ denotes a static spin-susceptibility, while $T_{\parallel}$ corresponds to the longitudinal relaxation time mentioned in the introduction above. The aim is to establish a direct comparison between Eqs. 7 and 14. For this purpose, let us use the first-order Taylor expansion of the KS spin-susceptibility [77]

$$\chi^{\text{KS}}_{\parallel}(\omega) \simeq \rho_F - i n_{e-h} \omega,$$  (15)

with $\rho_F = \rho_{F,\uparrow} + \rho_{F,\downarrow}$ the DOS at the Fermi level and $n_{e-h} = \pi(\rho_{F,\uparrow}^2 + \rho_{F,\downarrow}^2)/2$ the density of electron-hole excitations of the same spin channel. By inserting $\chi^{\text{KS}}_{\parallel}(\omega)$ of Eq. 15 into Eq. 7, $\chi_{\parallel}(\omega)$ acquires a functional form in $\omega$ equal to that of $\chi^{\text{Bl}}(\omega)$ in Eq. 14. This then allows to obtain an expression for the longitudinal relaxation time in terms of basic electronic properties (see Appendix A for details):

$$T_{\parallel} = \frac{U_{\parallel} n_{e-h}}{U_{\parallel} \rho_F - 1}.$$  (16)

The above expression shows that the longitudinal relaxation time is mostly settled by the magnitude of electron-hole excitations weighted by the XC kernel (see the numerator of Eq. 16); this product is of order $1/eV$, hence settling the overall time scale of $T_{\parallel}$ as fs. Furthermore, Eq. 16 shows that $T_{\parallel}$ diverges as $U_{\parallel} \rho_F \rightarrow 1$ (see the unitless denominator in the equation) and hence its magnitude can be strongly modified in this limit, i.e. as the system approaches the magnetic transition point. In the following, we first focus on quantitatively analyzing the ingredients of Eq. 16 and subsequently turn to $T_{\parallel}$ itself.

In order to compute reliable values for the kernel $U_{\parallel}$, one can make use of the static limit of Eq. 7, from which

$$U_{\parallel} = \rho_F^{-1} - \chi_{\parallel}^{-1}(0).$$  (17)
We note that $\chi(0)$ can be calculated by a standard ground state DFT calculation with a static magnetic field $\Delta B$ via $\chi(0) = \Delta m/\Delta B$, with $\Delta m$ the corresponding self-consistent change of the magnetic moment [97]. In Fig. 6 we show the calculated values of $\rho_F$ and $U_\parallel$ for several 3d and 4d adatoms deposited on Ag(100). The most important message conveyed by this figure is the large variation of $U_\parallel$ among different elements; while $U_\parallel \lesssim 0.5$ eV for most 4d elements, $U_\parallel \gtrsim 1.5$ eV for various 3d elements, reaching a maximum of one order of magnitude difference between Ru and Cr. A second important feature revealed by Fig. 6 is the distribution of $U_\parallel$ within each $d$-shell, whereby it is smallest at the ends of the row — case of Ti and Co among 3d, Nb and Ru among 4d — and highest in the middle of the row — case of Cr and Mn among 3d, Mo and Tc among 4d —, yielding an approximate inverted V-shape. We note that $\rho_F$ in Fig. 6 shows the opposite behavior, i.e. it is minimum for Cr and maximum for Co and Ru. This is consistent with Eq. 17, although we note strong deviations from the $U_\parallel \propto \rho_F^{-1}$ relationship (see in particular the case of Cr), revealing the importance of the term $\chi^{-1}(0)$ in Eq. 17.

The inverse of the static spin-susceptibility $\chi^{-1}(0)$ is closely connected to the magnetic equation of state, i.e. the dependence of the energy $E$ as a function of the magnetic moment $m$, via [97],

$$\chi^{-1}(0) = \frac{\partial^2 E(m)}{\partial m^2} \bigg|_{m=m_0} < 0. \tag{18}$$

In essence, the magnetic equation of state informs about how stable the magnetic solution is in comparison to the non-magnetic one. Fig. 7 shows the calculated $E(m)$ for the set of adatoms considered in Fig. 6 (calculations were carried out by employing DFT for a series of external magnetic fields [97]). This figure reveals that the energy difference between the non-magnetic and magnetic state,

$$\Delta E = E(m = 0) - E(m = m_0), \tag{19}$$

is of the order of eV and can largely vary for different adatoms. Importantly, the calculations show that 3d adatoms overall have a substantially larger $\Delta E$ than 4d adatoms. For Cr, for
Figure 7: Calculated energy as a function of the adatom’s magnetic moment. (a) and (b) 3d and 4d elements, respectively. DFT calculations are denoted by markers, while the solid lines are a fit to the calculations using \( E = \sum_{i=1}^{4} a_i m^{2i} \). The horizontal dashed lines mark the minimum energies for different adatoms.

instance, \( \Delta E \sim 3 \text{ eV} \), while for Ru \( \Delta E \sim 0.25 \text{ eV} \). Therefore, DFT predicts most 3d adatoms to be magnetically more stable than 4d ones, as expected. Furthermore, given that Eq. 18 together with Eq. 17 relates the XC kernel to the second derivative of the equation of state at \( M_0 \), one can establish an approximate connection between the depth of the minimum of \( E(M) \) and the value of \( U_\| \), as it is visible from the comparison of Figs. 7 and 6; the deeper the minimum, the larger \( U_\| \).

We note that following the above procedure, one can also extract the kernel in the non-magnetic ground state, i.e. the so-called Stoner XC parameter \( I_{xc} \) [98]. This can be achieved by considering the curvature of \( E(M) \) not at \( m = m_0 \) but at \( m = 0 \), as well as using the non-magnetic DOS in Eq. 17 instead of the magnetic one. As it is clearly visible from Fig. 7, the curvature is very different at \( m = 0 \) and \( m = m_0 \). Furthermore, \( \rho_F \) can also strongly vary from a magnetic to a non-magnetic calculation. As a consequence, the distribution of \( I_{xc} \) along the transition metal series first reported by Janak in Ref. [98] is very different to that of \( U_\| \) illustrated in Fig. 6.

Having analyzed the properties of \( \rho_F \) and \( U_\| \), we next focus on the longitudinal relaxation time \( T_\| \). The values calculated from Eq. 16 are plotted in Fig. 8(a) for 3d and 4d adatoms deposited on Ag(100) and Cu(111). \( T_\| \) is of the order of a few fs in all cases, being overall slightly larger for 4d than 3d adatoms, while the choice of substrate does not substantially affect it. Within each \( d \)-shell, \( T_\| \) is largest at the ends of the row while it is minimum for the half filled elements, thus resembling the behavior of \( \rho_F \) (compare Figs. 6 and 8(a)). Ru on Ag(100) has the highest value of \( T_\| \sim 50 \text{ fs} \), mainly as a consequence of the denominator of Eq. 16 being closer to zero than in other elements. In contrast, Cr and Mn have \( T_\| \sim 1 \text{ fs} \) in both substrates, i.e. nearly two orders of magnitude less than the aforementioned example. As a general feature, we note that the order of magnitude of \( T_\| \) is settled by the energy scale of the problem: all quantities involved in Eq. 16 are of the order of eV, whose corresponding time scale is fs. Therefore, the longitudinal relaxation of the spin considered here is extremely fast. The physical reason is the large exchange splitting dominating the relaxation process, which makes it energetically very expensive to modify the length of the moment due to the high energies involved.
Figure 8: Longitudinal relaxation time. (a) Calculated values for the longitudinal relaxation time $T_{∥}$ for magnetic $3d$ and $4d$ adatoms on Ag(100) (blue circles) and Cu(111) (red triangles). (b) Schematic illustration of two different processes contributing to the spin-relaxation time. Green and red dots respectively represent the excited and ground states. (b) Schematic description of the excitations contributing to $T_{∥}$ considered in this work. These take place in a continuum energy landscape of order eV and are driven by direct spin-conserving electron-hole transitions. (c) Illustration of the prototypical energy diagram used to describe the experiments of Refs. [41, 57, 22, 72]. The initially degenerate two ground states of maximum $S_z$ become non-degenerate by energy $\sim \) meV under externally applied magnetic fields of 1–10 T. Transitions between these two states then determine $T_1$, which take place via quantum tunneling.

4.1.1 Connection to experimental measurements

Let us next consider the experimental scenario regarding the measurement of the longitudinal spin-relaxation time. For this, we first note that in an experiment, several different mechanisms can contribute to this relaxation process, whose overall relaxation time is generally denoted as $T_1$. In this context, $T_{∥}$ calculated here is a particular contribution to $T_1$, which may include further contributions depending on the physical processes taking place. To the best of our knowledge, the first experimental technique that measures $T_1$ in magnetic single adatoms was developed by Loth and co-workers [41]. Within this STM-based technique, the spin-relaxation time was measured by monitoring the decay of electrons in excited states after the application of an all-electronic pump-probe scheme [41]. It is noteworthy that this scheme has so far only been applied to adatoms deposited on semi-insulating substrates, which are close to the atomic limit. The original work by Loth and co-workers measured $T_1 \sim 90$ ns for a Fe-Cu dimer on Cu$_2$Ni/Cu(100) [41]. A subsequent work by Rau and co-workers measured $T_1 \sim 200$ µs for a single Co atom on MgO/Ag(100) [57]. Lastly, Baumann and co-workers reported $T_1 \sim 90$ µs for a single Fe atom on MgO/Ag(100) [22], while in a recent work of Paul and co-workers on the same system, [72] the value of $T_1$ was enhanced up to the ms regime by fine-tuning external conditions such as the height of the STM tip. To conclude, we note that the reported time resolution of the measuring technique employed in the above experiments ranges between few ns to hundreds of ps.

All the above measured values of the spin-relaxation time are several orders of magnitude larger than the values of order fs that we discussed in this review for $T_{∥}$ (see Fig. 8(a)). Let us first
point out that all of the above experiments were performed under externally applied static magnetic fields that range between 1 T and 10 T. This, in turn, breaks the degeneracy of the spin ground states\cite{41}, a situation that is commonly modeled by a shifted discrete energy diagram as the one shown in Fig. 8(c). We note that excitations within such a diagram are not allowed to change the length of the spin moment (spin quantum number $S$ in this context), but only its projection (magnetic quantum number $S_z$). Therefore, the main spin-relaxation process contributing to $T_1$ within such an scheme involves transitions between the two non-degenerate states with same $S$ but opposite $S_z$ (see Fig. 8(c)). We note that their energy separation is of order meV, hence much smaller than the excitations of order eV involved in the change of the spin magnetic moment size considered for the calculation of $T_\parallel$, as schematically depicted in Fig. 8(b). Furthermore, given that direct transitions between the two non-degenerate states of Fig. 8(c) are virtually nonexistent, spin-relaxation in these conditions is driven by quantum tunneling processes, which are intrinsically much slower than the direct transitions considered here. These two considerations explain why the spin-relaxation time measured under the mentioned experimental conditions is several orders of magnitude larger than the values of $T_\parallel$ analyzed in this review.

It is apparent that, in order to experimentally access the dynamics encoded into $T_\parallel$, a measuring scheme based on ultrafast techniques that modify the length of the spin magnetic moment is required. Considering the technological developments within STM measuring techniques\cite{99,100,101}, accessing the fs time scale of magnetic adatoms seems to be a reasonable goal for the near future by, e.g. using ultrafast laser pulses, a breakthrough that would allow to monitor the ultrafast spin-dynamics analyzed here.

### 4.2 Transverse component

While longitudinal spin-susceptibility contains excitations that modify the spin density, the transverse one describes damped precessional motion of the spin moment\cite{96}. This motion, in turn, is described by the imaginary part of the enhanced spin-susceptibility, $\text{Im}\chi_\pm(\omega)$, a quantity that has been analyzed in depth in Sec. 3 (in particular, see Fig. 2). The characteristic form of the spin-excitation hosted by $\text{Im}\chi_\pm(\omega)$ and its connection to the transverse spin-relaxation time is illustrated in Fig. 9(a). As discussed in Sec. 3, $\text{Im}\chi_\pm(\omega)$ is mainly determined by the resonance frequency, $\omega_{\text{res}}$, which is settled by the SOC, and the width of the spin-excitation, $\Gamma$, which is proportional to the damping term $\eta$ (see in particular Eq. 52 in Appendix B). Noteworthily, the main contribution of the hybridization to $\Gamma$ is proportional to the electron-hole excitations of opposite spin channel, $n'_{e-h} = \pi \rho_{F,\uparrow} \rho_{F,\downarrow}$\cite{80}:

$$\Gamma \approx \frac{n'_{e-h}}{Q} \omega_{\text{res}}$$

with $Q = \text{Re} \frac{\partial \chi_{\pm}^{KS}(\omega)}{\partial \omega}|_{\omega=0}$. We note that the order of magnitude of $\omega_{\text{res}}$ ranges between $10^{-2} - 1$ meV while $n'_{e-h}/Q$ is a unitless fraction that is typically of order unity.

Importantly, a finite width corresponds to a finite damping of the precessing magnetic moment and is thus directly linked to the transverse spin-relaxation time (see Appendix B and Eq. 20):

$$T_\perp = \frac{2}{\Gamma} \propto (n'_{e-h})^{-1}.$$
Figure 9: Transverse relaxation time. (a) Schematic illustration of a transverse spin-excitation of a single adatom contained in Im $\chi_{\pm}(\omega)$. The resonance frequency, $\omega_{\text{res}}$, and the width, $\Gamma$, are indicated in the figure. (b) Calculated values for the transverse relaxation time $T_{\perp}$ for magnetic 3$d$ and 4$d$ adatoms on Ag(100) (blue circles) and Cu(111) (red triangles). Note the logarithmic scale in the y axis.

We note that, while $T_{\parallel}$ in Eq. 16 is directly proportional to the density of spin-conserving electron-hole excitations $n_{e-h}$, $T_{\perp}$ above is inversely proportional to the spin-flip counterpart $n'_{e-h}$. Let us also note that an expression with similar physical implications has been derived in the context of spin models with Kondo exchange in open quantum systems, the so-called non-adiabatic contribution to the spin-decoherence rate [23] (see in particular Eq. (70) in that reference).

In Fig. 9(b) we show calculated values for $T_{\perp}$ extracted from the corresponding spin-excitation width (see Fig. 2). We note that the variation of $T_{\perp}$ among adatoms is considerably larger (up to two orders more) than that of $T_{\parallel}$, shown in Fig. 8(a). This is a consequence of the large variation of the width $\Gamma$ of atomic spin-excitations, which can range from $10^{-2}$ meV to few meV [53, 81], i.e. nearly three orders of magnitude change. This, in turn, can be linked to the electronic DOS at the Fermi level via Eq. 20; adatoms where the DOS peak of the d-states lies close to the Fermi level, case of Ti, V, Fe and Co in Fig. 1(c), tend to be much more hybridized than those where only the tail of the DOS peak lies at the Fermi level, case of Cr and Mn in Fig. 1(c). In this way, Cr and Mn acquire large relaxation times of $T_{\perp} \sim 10^{1}-10^{2}$ ps, while strongly hybridized adatoms such as Co, Nb and Ru have $T_{\perp} \sim 10^{-1}-10^{-2}$ ps. In all cases, the transverse spin-relaxation process is orders of magnitude slower than the longitudinal one, $T_{\perp} \gg T_{\parallel}$.

To conclude, let us note that the trend of $T_{\perp}$ within each d-shell row is opposite to that shown by $T_{\parallel}$ (see Fig. 8(a)). This comes as a consequence of the dependence on the density of electron-hole excitations, with $T_{\parallel} \propto n_{e-h}$ (see Eq. 16) and $T_{\perp} \propto (n'_{e-h})^{-1}$ (see Eq. 20). In fact, $T_{\parallel}$ and $T_{\perp}$ can be formally related to each other considering the relation between the spin-conserving and spin-flip electron-hole contributions, namely $\rho_{F}^{2}/2 = n_{e-h} + n'_{e-h}$. From Eq. 16 for $T_{\parallel}$ and Eqs. 20 and 21 for $T_{\perp}$ one can then infer the following expression:

$$\frac{\rho_{F}^{2}}{2} \simeq T_{\parallel} \frac{U_{\parallel}}{U_{||}} - 1 + 2 \frac{Q}{T_{\perp}} \frac{\omega_{\text{res}}}{\omega_{\text{res}}}.$$  \hspace{1cm} (22)

The unifying concept behind the above relationship between $T_{\parallel}$ and $T_{\perp}$ is the hybridization of substrate electrons with the d-states of the transition metal adatoms, which in essence gives rise
to a finite total $\rho_F$. However, despite the formal relationship, the fact that $T_\parallel$ and $T_\perp$ in Eq. 22 have fundamentally different prefactors makes the time scale of the two relaxation constants differ by nearly three orders of magnitude.

4.2.1 Connection to experimental measurements

Next, we consider several experimental measurements of spin-excitation lifetimes of different single adatoms and connect them to this work. The lifetime of an atomic spin-excitation can be experimentally accessed from the width of the step observed in IETS $dI/dV$ measurements, which provides a measure of $\Gamma$. Given that the energy resolution of this technique is $10^{-1}$ meV at best [9], the longest lifetimes that can be inferred following this procedure are of order 10 ps (see Eq. 21). This type of experiments can measure adatoms deposited on both metallic and semi-insulating substrates; as a general trend, the latter induce a larger lifetime than the former due to a far smaller electronic hybridization. We begin by considering Ref. [8], where Khajetoorians and co-workers estimate the spin-excitation lifetime of a Fe adatom deposited on metallic Cu(111) to be 0.2 ps, in very good quantitative agreement with the calculated value $T_\perp = 0.15$ ps for the same system (see Fig. 9(b)). Noteworthily, when the same atom is deposited on metallic Pt(111), the measured lifetime is increased by more than an order of magnitude [9, 14]. Note that one finds a similar variation between the two substrates considered here for the elements Ti, V, Cr and Mo, as it can be checked in Fig. 9(b). Focusing next on the semi-insulating Cu$_2$Ni/Cu(100) substrate, a lower bound of $\sim 10$ ps has been experimentally estimated for Fe [3, 71], Mn [71] and Co [40] adatoms, although it is possible that the actual lifetimes are substantially larger. In fact, the calculations on Cr and Mn, which are the elements with smallest hybridization and thus the ones closest to the semi-insulating limit, show that $T_\perp$ can reach up to $10^3$ ps (see Fig. 9(b)); hence, it is not unlikely that the lifetimes of the aforementioned adatoms on Cu$_2$Ni/Cu(100) could be of the same order of magnitude. Last, it is worth noting the case of Co on MgO [57], which, despite being a semi-insulating substrate, yields a relatively short spin-excitation lifetime of $\sim 0.5$ ps, i.e. a common value for adatoms deposited in metallic substrates analyzed in this review (see Fig. 9(b)).

5 Paramagnetic spin-excitations

In this last section, we switch the main subject of study from magnetic to nonmagnetic adatoms, i.e., adatoms where the Stoner criterion for magnetism is nearly fulfilled. In particular, we analyze their dynamical properties, which can exhibit fingerprints of magnetism in the form of well-defined features in the spin-excitation spectrum, i.e., paramagnetic spin-excitations (PSE). Noteworthily, these excitations can be viewed as persistent spin-fluctuation modes that are activated by temperature and thus produce a measurable impact on properties such as specific heat or electron effective-mass enhancement [73, 67]. Thus, the same concept that leads to the destabilization of the magnetic moment seen in Sec. 3 can in this case instead be used to generate a signal of magnetic origin.

As in previous sections, the central quantity for the discussion is the spin-excitation spectrum,
which in this case is encoded into the paramagnetic spin-susceptibility (see Eq. 7): [69]

\[
\chi(\omega) = \frac{\chi_{KS}(\omega)}{1 - I_s \chi_{KS}(\omega)}. \tag{23}
\]

Above, \(I_s\) denotes the so-called Stoner parameter, which plays the role of the exchange-correlation kernel in the adiabatic local spin-density approximation. It is noteworthy that the static limit of Eq. 23 recovers a local form of the familiar Stoner theory that provides the well-known criterion for magnetism, i.e.

\[
\frac{\partial^2 E(m)}{\partial m^2} \bigg|_{m=0} = \chi(0)^{-1} < 0 \Rightarrow I_s \rho_F > 1, \tag{24}
\]

where we used \(\chi_{KS}(0) = \rho_F\) [69]. In essence, the product \(I_s \rho_F\) quantifies the competition between the exchange interaction, which enhances the tendency towards magnetism of electrons in localized orbitals, and substrate hybridization, which induces delocalization of the adatom’s electrons and therefore acts against magnetism, thus playing the role of the kinetic energy in the standard Stoner theory. It is interesting to note that even if an adatom does not fulfill the Stoner criterion, it can still develop dynamical PSE provided the details of the electronic structure make the denominator of Eq. 23 vanishingly small at a finite frequency.

### 5.1 Searching for the optimal candidates via the Stoner criterion

We begin by characterizing the set of 3d, 4d and 5d transition metal adatoms that could potentially exhibit PSE. For this purpose, we consider the Stoner criterion of Eq. 24 and search for adatoms whose Stoner product is close to 1. In Fig. 10(a) we plot \(I_s \rho_F\) for several adatoms deposited on the Ag(100) and Cu(111) substrates analyzed throughout the review. In addition, we have also included the MgO/Ag(100) substrate in order to analyze the differences introduced by a semi-insulating layer. We first note that the Stoner products of all the adatoms are below or slightly above 1. As a general trend, the calculations show that the metallic substrates Ag(100) and Cu(111) host adatoms whose Stoner product is closer to the critical value 1 as compared to semi-insulating MgO/Ag(100). This is mainly due to the small \(\rho_F\) in the latter, as tabulated in Table 2. Among the two metallic substrates, Ag(100) hosts adatoms whose Stoner product

![Figure 10: Ground state electronic properties of various 3d, 4d and 5d transition metal adatoms.](image)

- (a) Calculated Stoner product for adatoms deposited on Ag(100) (squares), Cu(111) (circles) and MgO/Ag(100) (triangles).
- (b) Calculated non-magnetic DOS of adatoms on Ag(100). The vertical line denotes the Fermi level.

\[
\left(\frac{\partial^2 E(m)}{\partial m^2}\right)_{m=0} = \chi(0)^{-1} < 0 \Rightarrow I_s \rho_F > 1,
\]

where we used \(\chi_{KS}(0) = \rho_F\) [69]. In essence, the product \(I_s \rho_F\) quantifies the competition between the exchange interaction, which enhances the tendency towards magnetism of electrons in localized orbitals, and substrate hybridization, which induces delocalization of the adatom’s electrons and therefore acts against magnetism, thus playing the role of the kinetic energy in the standard Stoner theory. It is interesting to note that even if an adatom does not fulfill the Stoner criterion, it can still develop dynamical PSE provided the details of the electronic structure make the denominator of Eq. 23 vanishingly small at a finite frequency.
<table>
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<tr>
<th></th>
<th>Sc</th>
<th>Ni</th>
<th>Cu</th>
<th>Y</th>
<th>Rh</th>
<th>Pd</th>
<th>Ir</th>
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<td>3.51</td>
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<td>0.32</td>
<td>0.28</td>
<td>0.27</td>
<td>0.39</td>
<td>0.40</td>
<td>0.36</td>
</tr>
<tr>
<td>$I_s \rho_F$</td>
<td>1.09</td>
<td>0.88</td>
<td>0.11</td>
<td>0.38</td>
<td>0.95</td>
<td>0.33</td>
<td>1.12</td>
<td>0.32</td>
</tr>
</tbody>
</table>

Table 2: Calculated values for the DOS at the Fermi level ($\rho_F$), Stoner parameter ($I_s$) and Stoner product ($I_s \rho_F$) for several 3d, 4d and 5d adatoms on Ag(100).

**Figure 11:** Spin-excitation spectrum of selected adatoms. (a) Solid lines illustrate the calculated density of PSE as given by $\text{Im} \chi(\omega)$ (Eq. 23) of selected 3d, 4d and 5d transition metal adatoms deposited on the metallic Ag(100) substrate. Dashed lines denote the approximation of Eq. 25. Note that both Eqs. 23 and 25 give rise to PSE. (b) and (c) Density of PSE as given by $\text{Im} \chi(\omega)$ for Rh and Ni adatoms, respectively, for magnetic fields of up to 18 T (both figures share the same legend). (d) Same as in (c) but for larger magnetic fields of up to $10^3$ T. The inset depicts the evolution of the PSE’s amplitude (see Eq. 26) as a function of the magnetic field.

are closest to 1, with $I_s \rho_F$ ranging between $\sim [1 - 0.1, 1 + 0.1]$ for Sc, Ir, Rh and Ni adatoms. Therefore, in the rest of this Section we focus on discussing the Ag(100) substrate in detail, as it illustrates best the concept of PSE.

In Fig. 10(b) we show the calculated DOS of the considered 3d, 4d and 5d adatoms. The figure shows that the large peaks coming from localized $d$-states acquire a finite width, which ranges from $\sim 0.2$ eV in the case of Cu, up to $\sim 1$ eV in the case of Sc and Y. This feature reveals a strong hybridization of the adatom’s $d$-states with the electron bath of the substrate [52, 80]. As a consequence, the DOS acquires a finite value at the Fermi level, $\rho_F$, which is listed in Table 2 for all adatoms, alongside with the calculated Stoner parameter and the resulting Stoner product.

We note that the largest $\rho_F$ corresponds to Rh due to the large width of the nearly filled $d$-peak in the DOS and its proximity to the Fermi level (see Fig. 10(b)). It is also noteworthy that Sc and Ir, which are predicted to be slightly magnetic, have a smaller $\rho_F$ than Rh and Ni. This is compensated by the Stoner parameter, which is almost 50% larger as compared to Ni and Rh. In this respect, the case of Ni is specially interesting, given that the calculated $I_s$ for the adatom is almost 50% smaller compared to the bulk value [98]. This drop of $I_s$ is the reason why Ni is magnetic in bulk but non-magnetic as an adatom on Ag(100) and other metallic substrates [102, 103].
5.2 Fingerprints of magnetism in the dynamical regime

In Fig. 11(a) we illustrate the calculated spin-excitation spectra as given by \( \text{Im} \chi(\omega) \) from Eq. 23, where all calculations were done considering the non-magnetic ground state. Interestingly, Fig. 11(a) reveals peak-like structures resonating at frequencies below 100 meV for Rh, Ni, Ir and Sc adatoms. This is exceptional, as most non-magnetic elements exhibit a featureless spectrum owing to a complete overdamping of the excitations. Rh represents the most favorable case, displaying a well-defined peak at \( \omega_{\text{res}} \sim 20 \text{ meV} \) and a width of \( \Delta \sim 50 \text{ meV} \), the associated lifetime being \( \sim 30 \text{ fs} \). It is noteworthy that both the lifetime and the height of the peak, the latter being related to the intensity of the excitation, are only one order of magnitude smaller than those of usual transverse spin-excitations of magnetic adatoms studied in Sec. 3 (see in particular Fig. 2) and that are accessible by IETS (see, e.g., Fe on Cu(111) in Refs. [8, 53]). On the other extreme, Pd in Fig. 11(a) shows a highly overdamped resonance at around 600 meV (see figure inset) whose intensity is an order of magnitude smaller than that of Rh. Therefore, the calculations reveal the existence of PSE whose resonance frequency and width vary strongly depending on the adatom.

Next, we focus on characterizing the physical mechanism behind PSE that allows an interpretation of the \textit{ab initio} results displayed in Fig. 11(a). For this purpose, let us consider the frequency expansion of the paramagnetic KS spin response function up to linear order, i.e., \( \chi^{\text{KS}}(\omega) = \rho_F + i \alpha \omega + O(\omega^2) \). One can show [77] that the linear expansion coefficient is well approximated by \( \alpha \sim -\pi \rho_F^2 / 4 \). Therefore, the spin-excitation spectrum within this approximation is given by a simple expression involving only the DOS at \( E_F \) and the Stoner parameter:

\[
\text{Im} \chi(\omega) = \frac{\pi}{4} \frac{\rho_F^2 \omega}{(1 - I_s \rho_F)^2 + (\frac{\pi}{4} I_s \rho_F^2 \omega)^2}.
\]

The above expression can be readily computed by using the calculated values for \( \rho_F \) and \( I_s \) shown in Table 2: the results are displayed in Fig. 11(a) (see dashed lines). A comparison to the full \textit{ab initio} calculations (solid lines) reveals a very good agreement for frequencies below 100 meV in the case of Rh, Ir and Sc, where both the peak and width are properly described within \( \lesssim 10\% \) relative error. This error is considerably larger in the case of Ni, indicating the importance of higher order expansion terms in \( \omega \) for this case. Finally, the peak for Pd is far beyond the limit of small frequencies and therefore the approximation of Eq. 25 breaks down.

Proving Eq. 25 to be an accurate approximation of the full spin-excitation density given by Eq. 23 is extremely convenient, as the former provides an analytical interpretation for the origin of PSE in terms of just \( \rho_F \) and \( I_s \), two basic electronic properties of adatoms. Indeed, the resonance frequency, linewidth and amplitude of PSE predicted by Eq. 25 can be cast into simple expressions:

\[
\omega_{\text{res}} = \frac{4}{\pi} \frac{1 - I_s \rho_F}{I_s \rho_F^2}, \quad \Delta = 2\sqrt{3} \omega_{\text{res}},
\]

\[
A \equiv \text{Im} \chi(\omega_{\text{res}}) = \frac{1}{2I_s \rho_F |1 - I_s \rho_F|}.
\]

Interestingly, a potential measurement of the above quantities would directly yield experimental estimates for \( \rho_F \) and \( I_s \). In closer inspection, one recognizes the Stoner product \( I_s \rho_F \) as the key quantity in Eq. 26; as \( I_s \rho_F \to 1 \) (i.e., magnetic instability), the resonance frequency as
5.3 Potential means of measurement: connection to dI/dV curves

We next focus on assessing the potential impact of PSE on the dI/dV signal as measured in IETS experiments, the technique of choice for measuring magnetic excitations (see, e.g., Refs. [8, 9, 105]). The corresponding minimal setup is illustrated in Fig. 12, which displays an STM tip measuring the adatom’s excitations under an applied external magnetic field, denoted as $B$. We first notice that PSE respond to magnetic fields by shifting their resonance frequency. This is quantitatively demonstrated in Figs. 11(b) and 11(c), where the calculated spin-excitation spectra are shown for Rh and Ni adatoms, respectively, under $B$ fields of $\sim 10$ T that are achievable in state-of-the-art laboratories (see, e.g., Refs. [9, 57, 38]). Noteworthily, while the PSE of Rh shifts towards larger frequencies as $B$ is increased (see Fig. 11(b)), the PSE of Ni exhibits the opposite behavior (see Fig. 11(c)). This difference arises from the fact that magnetic fields induce an effective modification of Stoner product, i.e., $I_s \rho_F \rightarrow \xi(B) I_s \rho_F$, where $\xi(B)$ is a term that depends both on the magnetic field as well as on the adatom’s electronic structure [77]. In particular, the details of the latter make $\xi(B) > 1$ for Ni while $\xi(B) < 1$ for Rh [77], leading to the aforementioned divergent responses in accordance with Eq. 26.

Remarkably, when strong enough magnetic fields are applied to Ni, the modified Stoner criterion...
can be tuned towards the critical point, as shown in Fig. 11(d). As a consequence, the PSE’s resonance frequency approaches the origin in a singular way while the amplitude of the excitation is enhanced by as much as two orders of magnitude for $B \sim 500$ T. It is interesting to note that this critical behavior is also present on the $B$-field dependence of the induced magnetic moment $m$, as shown in the inset of Fig. 12. While Rh shows a continuous dependence, Ni reveals a discontinous transition at approximately the critical field value $B \sim 500$ T, above which the system enters a magnetic regime where the internal exchange field effectively contributes to $m$ on top of the external Zeeman field, featuring the atomic version of a quantum phase transition.

We note that, although such large $B$ fields are clearly out of reach for current experiments, this feature could be potentially observed, e.g., via the proximity effect, by placing a magnetic adatom in the neighborhood of the non-magnetic one (see Fig. 12). Calculations show that the former can induce on the latter a magnetic moment of the same order of magnitude as the one induced by the fields of Fig. 11(d) [104], thus mimicking the action of large magnetic fields.

Next we evaluate the impact of PSE on the $dI/dV$ signal of an IETS measurement. For such purpose we consider the so-called Tersoff-Hamann approximation [106, 107], which relates the tunneling conductance to the electronic DOS at the tip position, which is renormalized by the adatom’s excitations. The latter quantity can be accessed by means of a technique that combines many-body perturbation theory with the TDDFT scheme; details can be found in Ref. [54]. The central object within this formalism is the electron self-energy, $\Sigma$, which contains the interactions between the tunneling electrons from the tip at bias voltage $V$ and the adatom’s PSE. It is particularly revealing to inspect its imaginary part [54],

$$\text{Im} \Sigma(V_F) = -\int_0^V d\omega \rho(V_F + \omega) \text{Im} \chi(\omega),$$

with $\rho(E)$ the energy-dependent DOS, $V_F = E_F + V$ and $E_F$ the Fermi energy. The calculated Im $\Sigma(V_F)$ is shown in Fig. 13(a) for Rh under various magnetic fields of up to 18 T. The results reveal a clear step for positive bias voltage that saturates at $\sim 100$ meV, i.e., after the PSE peak has been integrated (see Eq. 27). Note also that the calculated self-energy slightly varies as a function of the magnetic field. When larger magnetic fields are applied, as illustrated in
Fig. 13(b) for the case of Ni, the critical behavior of the PSE (see Fig. 11(d)) translates into a clear maximum at the value of the critical field, where $\text{Im} \Sigma(V_F)$ increases by an order of magnitude.

The presence of PSE has a broad effect on the renormalization of the DOS at the vacuum, where IETS tips measure the signal. In particular, the energy derivative of the renormalized DOS (rDOS) is a quantity that is linked to the $d^2I/dV^2$ curve measured by IETS [54]. The former quantity is displayed in Fig. 13(c) for Rh, where the magnetic field dependence is clearly visible. Noteworthily, the calculations demonstrate that the tunneling electrons from the tip are able to trigger the PSE, leading to a peak in the meV region that, furthermore, reacts to external magnetic fields by shifting its resonance frequency as well as substantially modifying its intensity. We also note the strong asymmetric distribution between positive and negative frequencies, a feature that emerges from the background electronic structure [54] and is commonly present in $d^2I/dV^2$ curves measured on magnetic adatoms (see, e.g., Refs. [1, 8, 108, 9, 109]). On the other hand, when Ni is driven into the critical regime as in Fig. 13(d), the calculations reveal a huge change of the signal’s intensity as the PSE approaches the critical point. This analysis therefore shows that magnetism offers a prime way of manipulating PSE, enabling to discern them from other excitations of similar energy but non-magnetic origin, such as phonons.

6 Outlook

In this review, we have presented a state-of-the-art treatment based on TDDFT of several dynamical aspects of single-adatoms deposited on metallic substrates. As the first main topic, we have shown that the magnetic stability defined by the MAE can be severely reduced as a consequence of quantum spin-fluctuations that arise mainly from Stoner spin-flip excitations. Our analysis has quantitatively shown how protection against fluctuations is gained by reducing the hybridization with substrate electrons as well as maximizing the adatom’s local magnetic moment. In this context, future work should address an important contribution that has not been considered here, namely the vibrational contribution of phonons, which is believed to be key in systems of great importance like Ho on MgO/Ag(100) [38]. In particular, a reliable ab initio scheme for the calculation of the electron-phonon coupling [110, 111] and its effect on the magnetic stability of single-adatoms is necessary in order to complement the knowledge gained on the electron-electron side. A second important ingredient that should be further considered in the future from an ab initio viewpoint is the destabilizing effect of external particles, such as the tunneling electrons coming from an STM tip or X-ray photons of an XMCD measurement. In both cases, it is experimentally clear that the magnetic stability and spin lifetimes are severely affected by the intensity or flux of the incoming particles (see e.g., Refs [38, 42, 72].) First steps towards incorporating such effects into an ab initio scheme have been already taken in Ref. [54]: pending issues include the incorporation of SOC into the formalism as well as establishing a direct connection between the renormalization induced by the tunneling electrons and the magnetic stability of the adatom.

The conclusions drawn from the single-adatom case serve also as a solid platform for tackling more complex nanomagnets composed of several exchange-coupled magnetic atoms. This play-
ground is particularly attractive given that several STM experiments have reported clear observations of stable magnetic moments in this type of composite system (see e.g., Refs. [112, 63, 113]). This strongly suggests that local fluctuation effects decrease with increasing cluster size, a feature that is in qualitative accordance with one main finding presented throughout the review, namely that the relative impact of the fluctuations becomes smaller the larger the magnetic moment gets. This general trend, however, is likely to be modified by characteristics associated to clusters, such as the appearance of optical modes in the spin-excitation spectrum (i.e. additional peaks in $\text{Im } \chi_\perp(\omega)$, see e.g., Refs. [7, 52]) that would contribute to the fluctuations as seen from Eq. 6. On top of this, the impact of the fluctuations on the magnetic interactions among different adatoms remains to be fully explored and clarified.

Throughout the review we have also paid close attention to the time-scales of different spin-relaxation processes. We have studied two particular contributions, namely the longitudinal and transversal spin-relaxation times, providing their connection to the electronic structure of single-adatoms and, when possible, comparing the $ab\ initio$ data to experimental measurements. In the case of $T_\perp$, we have shown that it is governed by atomic spin-excitations that induce the precessional motion of the magnetic moment, hence this relaxation time contributes to $T_2$ as commonly defined in the standard literature [96, 114]. Interestingly, there is yet another contribution to $T_2$ that has recently attracted a great interest in the single-adatom community, namely the adiabatic spin-decoherence time (see e.g., the review [23]). In fact, very recent EPR-based experiments have provided hard numbers for this quantity [22, 115], which is key for the potential use of single-adatoms as quantum devices. The $ab\ initio$ calculation of this quantity is a challenging task for the future due to its pure quantum origin, which is not trivial to capture within the mean-field approach of DFT. Regarding the longitudinal spin-relaxation time, we have argued that, although currently available techniques cannot monitor the femtosecond regime of magnetic single adatoms, it is reasonable that this can be achieved in the near future, e.g., by employing STM-integrated ultrafast laser schemes [99, 100, 101], thus giving access to the ultrafast spin-dynamics described in this work.

As the last main topic, we have analyzed paramagnetic spin-excitations, i.e. fingerprints of magnetism in nonmagnetic adatoms, as well as means of detecting them in IETS experiments. Remarkably, a very recent measurement [116] of the conductance associated to a single Pd adatom deposited on Pd(111) has been interpreted as being strongly affected by paramagnon scattering, i.e., a potential indirect experimental observation of PSE. This opens up a promising range of applications for non-magnetic adatoms in nanotechnology, which encodes and manipulates information into excitation modes like PSE.

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Appendices

A Longitudinal Bloch equation

The Bloch equation for the longitudinal change of the magnetization under the effect of a time-dependent perturbation $H_1(t)$ along the longitudinal direction can be written as [96]

$$\frac{d m_z(t)}{dt} = \chi^{\text{Bl}}_0 H_1(t) - m_z(t) T_\parallel,$$

with $\chi^{\text{Bl}}_0$ a static spin-susceptibility. The above equation describes how $m_z(t)$ comes back to equilibrium with a characteristic relaxation time $T_\parallel$ after being perturbed by $H_1(t)$. Using $f(t) = \int \omega f(\omega)e^{-i\omega t}$ for both $m_z(t)$ and $H_1(t)$ we can write Eq. 28 in the frequency domain,

$$m_z(\omega) \left(-i\omega T_\parallel + 1\right) = \chi_0 H_1(\omega) \Rightarrow \frac{m_z(\omega)}{H_1(\omega)} = \chi^{\text{Bl}}(\omega) = \frac{\chi^{\text{Bl}}_0}{1 - i\omega T_\parallel},$$

where $\chi^{\text{Bl}}(\omega)$ is the enhanced spin-susceptibility. The real and imaginary parts of the above equation read

$$\text{Re} \chi^{\text{Bl}}(\omega) = \frac{\chi^{\text{Bl}}_0}{1 + (\omega T_\parallel)^2},$$

$$\text{Im} \chi^{\text{Bl}}(\omega) = \frac{\chi^{\text{Bl}}_0 \omega T_\parallel}{1 + (\omega T_\parallel)^2}.$$

Next, we consider the Taylor expansion of the KS spin-susceptibility (see Eq. 15 in the main text),

$$\chi^{\text{KS}}_\parallel(\omega) \simeq \rho_F - i n_{-h}\omega.$$ 

The first-order expansion coefficient $n_{-h} = \pi(\rho^2_F \uparrow + \rho^2_F \downarrow)/2$ has been calculated in the Supplemental Material of Ref. [77]. Inserting $\chi^{\text{KS}}_\parallel(\omega)$ of Eq. 32 into the definition of the TDDFT enhanced spin-susceptibility $\chi(\omega)$ (see Eq. 7 of the main text), the imaginary part $\text{Im} \chi_\parallel(\omega)$ can be cast in the following way,

$$\text{Im} \chi_\parallel(\omega) = \frac{n_{-h}\omega}{(1 - U_\parallel \rho_F)^2} \frac{1}{1 + \left(\frac{U_\parallel n_{-h}\omega}{1 - U_\parallel \rho_F}\right)^2} \frac{1}{U_\parallel \rho_F} = \frac{\chi(0) \omega T_\parallel}{1 + (\omega T_\parallel)^2},$$

where $\chi(0) = \chi^{\text{Bl}}_0$.
where in the last step we used the expression for the static spin-susceptibility $\chi_\parallel(0) = \rho_F/(1 - U_\parallel \rho_F)$ and we defined

$$T_\parallel = \frac{U_\parallel n_{e-h}}{1 - U_\parallel \rho_F},$$

(34)

which is the result quoted in the main text in Eq. 16.

B Transverse relaxation within the Landau-Lifshitz-Gilbert equation

We consider the Landau-Lifshitz-Gilbert (LLG) equation describing the damped precessional motion of a magnetic moment placed in a static external magnetic field that has been perturbed by a time-dependent transverse magnetic field:

$$\frac{d\mathbf{m}}{dt} = -\gamma \mathbf{m} \times \mathbf{B}^{\text{ext}} + \eta \frac{m_0}{m_0} \times \frac{d\mathbf{m}}{dt},$$

(35)

with $\mathbf{B}^{\text{ext}} = B_0 \hat{z} + \mathbf{b}(t)$ and $\mathbf{b}(t) = b_x(t) \hat{x} + b_y(t) \hat{y}$. Here $B_0$ is the static part of the external field, and $\mathbf{b}(t)$ a small transverse perturbing field, $|\mathbf{b}(t)| \ll B_0$. The precession rate in Eq. 35 is set by $\gamma$, while the relaxation is controlled by $\eta$, namely the damping term. Writing $\mathbf{m}(t) = m_0 \hat{z} + m(t) \hat{x} + m_y(t) \hat{y}$, and linearizing the LLG equation yields the following equation of motion for the transverse components of the magnetization:

$$\frac{dm_x}{dt} = -\gamma B_0 m_y + \gamma m_0 b_y - \eta \frac{dm_y}{dt},$$

(36)

$$\frac{dm_y}{dt} = -\gamma m_0 b_x + \gamma B_0 m_x + \eta \frac{dm_x}{dt}.$$  

(37)

To illustrate the relaxation dynamics, consider a static external field ($\mathbf{b}(t) = 0$) and a small tilt of the magnetic moment at some initial time: $\mathbf{m}(t = 0) = m_0 (\cos \theta \hat{z} + \sin \theta \hat{x}) \approx m_0 \hat{z} + \Delta m \hat{x}$. Since the expected solution is a damped precession that relaxes towards the direction of the static magnetic field, we use the following ansatz corresponding to a circular precession that decays in time with a transverse relaxation time $T_\perp$:

$$m_x(t) = \Delta m e^{-t/T_\perp} \cos(\omega_0 t),$$

(38)

$$m_y(t) = \Delta m e^{-t/T_\perp} \sin(\omega_0 t).$$

(39)

Plugging the above ansatz back into the LLG Eqs. 36 and 37 we get

$$\frac{1}{T_\perp} \cos(\omega_0 t) + \omega_0 \sin(\omega_0 t) = \eta \omega_0 \cos(\omega_0 t) + \left(\gamma B_0 - \frac{\eta}{T_\perp}\right) \sin(\omega_0 t),$$

(40)

$$\frac{1}{T_\perp} \sin(\omega_0 t) + \omega_0 \cos(\omega_0 t) = -\eta \omega_0 \sin(\omega_0 t) + \left(\gamma B_0 - \frac{\eta}{T_\perp}\right) \sin(\omega_0 t).$$

(41)

The above equations can only be satisfied if the coefficients in front of the time-dependent sines and cosines match. We then have (both equations give the same pair of relations)

$$1 = \eta \omega_0,$$

(42)

$$\omega_0 = \frac{\gamma B_0}{1 + \eta^2}.$$  

(43)
Importantly, Eq. 42 shows that the transverse relaxation time is given by the product between the damping term $\eta$ and the characteristic frequency $\omega_0$.

We next turn to calculate the transverse dynamic spin-susceptibility within the LLG model. For this, we consider the following Fourier transforms,

$$b(t) = \int \frac{d\omega}{2\pi} e^{-i\omega t} b(\omega), \quad m(t) = \int \frac{d\omega}{2\pi} e^{-i\omega t} m(\omega).$$

(44)

Inserting the above expressions into the linearized equations 36 and 37 we obtain in frequency space

$$-i \omega m_x(\omega) = +\gamma m_0 b_y(\omega) + (i\eta\omega - \gamma B_0) m_y(\omega),$$

(45)

$$-i \omega m_y(\omega) = -\gamma m_0 b_x(\omega) + (\gamma B_0 - i\eta\omega) m_x(\omega).$$

(46)

The above can be simplified by considering the circular components $m_\pm = m_x \pm i m_y$, yielding

$$\omega m_\pm(\omega) = \gamma_\pm (m_0 b_\pm(\omega) - B_0 m_\pm(\omega)), \Rightarrow \Lambda_\pm(\omega) m_\pm(\omega) = b_\pm(\omega),$$

(47)

with $b_\pm = b_x \pm i b_y$, $\gamma_\pm = \pm \gamma/(1 \mp i \eta)$ and

$$\Lambda_\pm(\omega) = \frac{B_0}{m_0} + \frac{\omega}{\gamma_\pm m_0}. \quad (48)$$

It is apparent from Eq. 47 that the transverse spin-susceptibility can be obtained from the inverse of $\Lambda_\pm(\omega)$ defined above. After some algebra and picking the minus sign in Eq. 48 one obtains

$$\chi_{\text{LLG}}^\pm(\omega) = (\Lambda_-(\omega))^{-1} = \frac{m_0 \omega_0}{B_0} \cdot \frac{-\omega + (1 + \eta^2)\omega_0 + i\eta\omega}{(\omega - \omega_0)^2 + (\eta\omega_0)^2}. \quad (49)$$

The density of spin-excitations in the LLG model are thus described by a skewed Lorentzian in $\omega$:

$$\text{Im} \chi_{\text{LLG}}^\pm(\omega) = \frac{m_0 \omega_0}{B_0} \frac{\eta\omega}{(\omega - \omega_0)^2 + (\eta\omega_0)^2}. \quad (50)$$

The resonance frequency of the above function takes place at

$$\frac{d}{d\omega} \text{Im} \chi_{\text{LLG}}^\pm(\omega) = 0 \Rightarrow \omega_{\text{res}} = \sqrt{1 + \eta^2}\omega_0, \quad (51)$$

while the FWHM amounts to

$$\Gamma = 2\eta\omega_0 \frac{\sqrt{2 + 3\eta^2 + 2\sqrt{1 + \eta^2}}}{1 + \sqrt{1 + \eta^2}} \simeq 2\eta\omega_0. \quad (52)$$

We note that the above approximation is exact in the $\eta \to 0$ limit and involves only a $\approx 10\%$ relative error for $\eta = 1$, which is by far the maximum value that damping can get for single adatoms; for most of the elements analyzed in the main text we have $\eta \lesssim 0.5$ [81], so the approximation of Eq. 52 is indeed very good. Then, comparing Eq. 52 to Eq. 42 we arrive to the relation between the FWHM and the transverse relaxation time quoted in the main text:

$$T_\perp = 2\frac{\Gamma}{\Gamma}. \quad (53)$$
References


[104] For externally applied $B = 517$ T the magnetic moment of Ni is $m = 0.29 \mu_B$, while a Fe adatom placed as a second nearest neighbor induces a magnetic moment on Ni of $m = 0.51 \mu_B$.


