First-principles calculations of spin and angle-resolved resonant photoemission spectra of Cr(110) surfaces at the 2p - 3d resonance

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A first principles approach for spin and angle resolved resonant photoemission is developed within multiple scattering theory and applied to a Cr(110) surface at the 2p-3d resonance. The resonant photocurrent from this non ferromagnetic system is found to be strongly spin polarized by circularly polarized light, in agreement with experiments on antiferromagnetic and magnetically disordered systems. By comparing the antiferromagnetic and Pauli-paramagnetic phases of Cr, we explicitly show that the spin polarization of the photocurrent is independent of the existence of local magnetic moments, solving a long-standing debate on the origin of such polarization. New spin polarization effects are predicted for the paramagnetic phase even with unpolarized light, opening new directions for full mapping of spin interactions in macroscopically non magnetic or nanostructured systems.

In recent years, the theoretical description of absorption/photoemission spectroscopy in the X-ray region has been boosted by the merge of density functional theory (DFT) with many body approaches such as dynamical mean field theory [1, 2], many body perturbation theory [3–5] and by the development of time-dependent DFT [6]. However, second order processes, like resonant inelastic X-ray scattering (RIXS) and resonant photoemission (RPES), remain a major challenge for theory. For RPES, existing approaches are semiempirical [7–10], based on a well defined two-holes final state and on small clusters, and thus do not take into account the delocalization of intermediate states, the bandstructure of the system and multiple scattering effects in the propagation of photoelectrons.

The huge experimental output from RPES on correlated materials [7, 11–16] and the intriguing quest for a determination of local magnetic properties put forward by pioneering experiments [14–16] call for advancements in the theoretical description of this spectroscopy. In experiments on CuO and Ni, it was shown that the RPES photocurrent with circular polarized light is spin polarized in antiferromagnets [14, 15] and Curie paramagnets [16]. It was claimed that a specific combination of spin resolved spectra provides a direct measure of the local magnetic moments [14, 16]. The issue is of fundamental importance in the search for a tool to access the local magnetic properties in antiferromagnetic, magnetically disordered and/or nanostructured systems at their crossover with the transition temperature. The interpretation was however rejected on the basis of symmetry analysis [17], but explicit calculations predicting the line-shape and intensity of such fundamental signal are still lacking and remain highly desirable.

In this letter, we present the first ab-initio method for RPES in solids, based on a combined formulation within the real space multiple scattering (RSMS) approach [18, 19] and DFT, and its application to Cr(110) at the 2p-3d resonance. By comparing the antiferromagnetic (AFM) and Pauli-paramagnetic (PM) phase of Cr, we solve the long-standing debate about the possibility to determine local magnetic moments in macroscopically non magnetic systems by means of spin resolved RPES with circular polarized light. New interesting effects in the PM phase by unpolarized light suggest that other mechanisms are active and could be exploited for mapping the origin of the different spin polarization (SP) components in paramagnets and magnetically disordered systems.

Theoretical formulation. The cross section for valence band photoemission to a final state $|u, k\rangle$, where $u$ denotes a valence band hole and $k$ a photoelectron state, is given by

$$I(\omega, q, k) = \sum_v |T_{kv}(\omega, q)|^2 \delta(\epsilon_k - \epsilon_v - \hbar\omega)$$

where $\hbar\omega$ and $q$ are the photon energy and polarization. Here the independent particle approximation has been assumed (i.e., all many-electron eigenstates are single Slater determinants corresponding to the same effective one-electron hamiltonian). According to the Heisenberg-Kramers formula [20], the transition matrix element $T_{kv}(\omega, q)$ is the sum of a direct and a resonant term. In the latter, photon absorption leads to an intermediate state $|u, w\rangle$, with a core hole $|\bar{\xi}\rangle$ and an electron in a formerly unoccupied state $|u\rangle$, which decays to the final state $|u, k\rangle$ through a participator Auger process [20, 21]. To lowest order in the autoionization process, the transition matrix element is given by

$$T_{kv}(\omega, q) = \langle k|D_q|v\rangle + \sum_{cu} \frac{\langle kc|V(|u\rangle - |w\rangle\rangle}{\hbar\omega + \epsilon_c - \epsilon_u - i\Gamma} (u|D_q|c)$$

where $D_q$ is the dipole operator, $V$ the Coulomb operator and $\Gamma$ the width of the intermediate state. Speci-
tator Auger decay leads to different, namely two-hole fin-
al states and is not considered here. Participator and 
spectator channels can in principle be separated experi-
mentally by using a photon bandwidth smaller than the 
core-hole lifetime, as they show different photon energy 
dependence (linear for the participator, and no photon 
energy dependence for the spectator). Here we focus on 
the physical effects at the origin of spin polarization and 
dichroism as well as their directional-dependence in the 
“pure” participator channel.

The RPES intensity can be written in a compact form as

\[ I(\omega, q, k) = \sum_{ijLL} M_{iL}^{\sigma,q}(k) I_{jL'}^{ij}(\epsilon_v, \sigma) M_{jL'}^{\sigma,q}(k)^* \]

Here, \( i, j \) label atomic sites, \( L \equiv (lm) \) angular momentum and \( \sigma \) spin quantum numbers. \( \epsilon_v = \epsilon_k - \hbar \omega \) is the energy of the valence hole. The quantity \( I_{jL'}^{ij}(\epsilon_k \sigma_k, \epsilon_v \sigma_v) \) is the essentially imaginary part of the scattering path operator. It comes from the simplification of the sum over delocalized valence states through the so called optical theorem in RSMS \[22\] and it contains the bandstructure information. The matrix elements \( M_{iL}^{\sigma,q}(k) \) are given by

\[ M_{iL}^{\sigma,q}(k) = \sum_{jL'} B_{jL'}^{\sigma}(k) A_{jL',iL}(\epsilon_k \sigma_k, \epsilon_v \sigma_v) \]

The \( B_{jL'}^{\sigma}(k) \) are the key quantities in the RSMS approach and represent the multiple scattering amplitudes of the continuum state \( k \equiv (k \sigma_k) \). \[22\]. The matrix elements \( A_{jL',iL}(\epsilon_k \sigma_k, \epsilon_v \sigma_v) \) are given by the sum of the direct radiative process \( (A^D) \), the resonant process with direct Coulomb decay \( (A^C) \) and the resonant process with the exchange decay \( (A^X) \), see Eq. 1. \( A^D \) and \( A^C \) are site-
and spin-diagonal \((\sim \delta_{ij} \delta_{\sigma \sigma})\). We have

\[ A^D = \langle \epsilon_k L' \sigma | D_q | \epsilon_v u, L \sigma \rangle \]

\[ A^C = - \sum_{jU_L} \int_{E_F} \frac{I_{jL'}^{ij}(\epsilon_u \sigma_u)}{\hbar \omega + \epsilon_c - \epsilon_u - i \Gamma} \times \langle \epsilon_k L' \sigma, j' | V | \epsilon_v u, L \sigma, j' \epsilon_u L \sigma_u \rangle \langle j' \epsilon_u L \sigma_u | D | j' c \rangle \]

\[ A^X = \sum_{\epsilon_u L \sigma_u} \int_{E_F} \frac{I_{jL'}^{ij}(\epsilon_u \sigma_u)}{\hbar \omega + \epsilon_c - \epsilon_u - i \Gamma} \times \langle j \epsilon_k L \sigma_k, i c | V | j \epsilon_u L \sigma_u, k \epsilon_u L \sigma_u \rangle \langle \epsilon_u L \sigma_u | D | i c \rangle \]

The sums over unoccupied states \( u \) have been again simplified through the optical theorem. The exchange term \( A^X \) is not strictly site-diagonal because of the non-
locality of the exchange interaction together with the de-
localized nature of the states \( u \). In the RSMS approach the Coulomb matrix elements \( \langle k | V | u \rangle \) and \( \langle k | V | u \rangle \) can be exactly developed in one- and two-center terms. In metallic Cr, the Coulomb interaction is strongly screened. As a result, two-center terms are by at least one order of magnitude smaller than the one-center terms \[23\] and have been neglected here. In general, the 2p-3d excited intermediate states might display excitonic effects, which could be taken account for with a Bethe-Salpeter description [3, 5]. For Cr metal, these effects are quite small because of the large 3d band width \((\sim 7 \text{ eV})\) and efficient metallic screening of the core hole by nearly free \(4sp\) electrons, and thus neglected here.

Photoemission spectra from Cr(110) are calculated in 
RSMS with a cluster of 151 atoms (see Fig. 1a) and self-
consistent spin polarized potentials, obtained by a scalar 
relativistic LMTO \[24\] calculation for bulk Cr in the 
local spin density approximation. Except for the 2p core 
level, all states entering the RPES calculation are de-
veloped in RSMS. The 2p orbital is obtained by solv-
ing the scalar relativistic Schrödinger equation with self-
consistent spin-polarized LMTO potentials. The 2p\(3/2\) 
spin-orbit coupled states are then constructed using stan-
dard angular momentum algebra and the spin-orbit cou-
pling constant is taken from an atomic calculation \[25\].

We consider the AFM order of CsCl-type which is a 
good approximation to the true spin density wave (SDW) 
ground state of Cr. The calculated magnetic moment is 
0.74 \( \mu_B \) in reasonable agreement with experiment 
(0.62 \( \mu_B \)). At the \( (110) \) surface, the transverse SDW 
propagates along \([100]\) or \([010] \[26\]. Therefore, we take 
\( \epsilon_z = [001] \) as magnetization and spin-quantization axis throughout this paper. We also consider the Pauli PM 
state, corresponding to a non-magnetic calculation. Spin 
orbit (SO) coupling of the valence and continuum states is 
neglected (it is as small as 0.03 eV for Cr-3d \[27\]).

Results. The electronic structure of Cr(110) is well ac-
counted for in the RSMS approach as can be seen from 
the comparison between the local density of states (DOS) of a Cr atom in the cluster and of bulk Cr (Fig. 1b). Non-
resonant angle-resolved photoemission spectra (ARPES) 
are shown in Fig. 1. Differences with respect to exper-
iments \[28\] are expected as our approach does not con-
tain local many-body interactions and layer-dependent 
potentials, which could play a role for a quantitative de-
scription of the peak renormalization and dispersion be-
aviour of the energetic structures \[29\]. However, the 
main features of the experimental spectra are reproduced in 
the calculation, confirming that RSMS provides a reason-
able good description of valence band photoemission from 
metals as previously shown for Cu(111) \[22\].

Spin resolved, angle integrated PES and RPES spec-
tra are shown in Fig. 2 for the AFM phase and several 
photon energies across the \( L_3 \)-edge absorption threshold. 
Left circular polarized light incident along the magne-
tization axis \([001]\) is considered. In this “parallel” ge-
ometry the spectra, right polarized light produces the 
same spectra but with up and down spin exchanged. The 
maximum peak intensity as a function of photon energy is 
plotted in Fig. 2b and shows the expected Fano pro-
file. The first photon energy (551.0 eV) is too low to ex-
cite the core electron and so only direct PES is possible.
FIG. 1: (a) Cr(110) cluster used in the RSMS calculations. The two magnetic sublattices of the AFM state are in red and blue. (b) DOS in the AFM phase for a bulk atom (LMTO) and a central atom in the cluster (RSMS). (c) ARPES spectra from Cr(110) along the (001) azimuth for different polar angles \( \theta \) with respect to the surface normal. Unpolarized light along the [001] axis was considered. Experimental data from Refs [14, 16] with disordered and/or fluctuating magnetic moments, but a Pauli PM state, unambiguously proves that it is unrelated to local magnetic moments in non-ferromagnetic samples. The normal emission ARPES spectra (Fig 3a,b) (total spectra) for parallel geometry consist of a single peak at 0.8-0.9 eV binding energy, very similar to the low energy non-resonant spectrum in Fig.1c (0.8-0.9 eV), which reflects the small exchange splitting of the Fermi level into the unoccupied Cr 3d band. At \( h\nu = 585.1 \) eV, well above threshold, the resonant spectrum goes back to the non-resonant one.

The direct PES signal is non-spin-polarized as expected for the AFM phase. Appreciable spin-polarization is, however, found in RPES. This effect is here obtained for the first time through first-principles calculations, and confirms the experimental finding in CuO [14], that in AFM systems RPES at the 2p_{3/2}-3d resonance is spin-polarized when circular polarized light is used.

We now turn to angle and spin resolved spectra at maximum resonance (\( h\nu=554.4 \) eV), focusing on their four “fundamental” combinations (and their relation to local magnetic properties), constructed by different choices of photoelectron spin (\( \uparrow,\downarrow \)) and light helicity (+,−)=(left,right):

\[
\text{tot} \equiv (\uparrow+)+(\uparrow−)+(\downarrow+)+(\downarrow−) \quad \text{(total)} \\
\text{spr} \equiv (\uparrow+)+(\uparrow−)+(\downarrow+)+(\downarrow−) \quad \text{(spin-resolved)} \\
\text{dic} \equiv (\uparrow+)-(\uparrow−)+(\downarrow+)-(\downarrow−) \quad \text{(dichroic)} \\
\text{mix} \equiv (\uparrow+)-(\uparrow−)-(\downarrow+)+(\downarrow−) \quad \text{(mixed)}
\]

The “mixed” spectrum was the one considered in Refs [14, 16] and claimed to be sensitive to local magnetic moments in non-ferromagnetic samples.

When the photon energy is raised to 552.4 eV, just below the absorption edge, direct and resonant process interfere destructively, giving rise to the dip in the Fano profile. Strong resonant enhancement is observed between 552 and 554.5 eV (see e.g. the spectrum for 554.4 eV), which corresponds to transitions from the 2p_{3/2} level into the unoccupied Cr 3d band. At \( h\nu = 585.1 \) eV, well above threshold, the resonant spectrum goes back to the non-resonant one.

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The normal emission ARPES spectra (Fig 3a,b) (total spectra) for parallel geometry consist of a single peak at 0.8-0.9 eV binding energy, very similar to the low energy non-resonant spectrum in Fig.1c (\( \theta = 0^\circ \)). AFM and PM spectra are almost identical except for a small shift of \( \sim 0.1 \) eV, which reflects the small exchange splitting of the AFM Cr-3d bands. The dichroic (dic) and spin-resolved (spr) signals vanish for both PM and AFM phase, as expected since the system is globally non-magnetic in both cases, and the set up is non chiral.

However, the mixed signal is non-zero with a large amplitude (\( \sim 1/3 \) of total), in agreement with the experimental results in AFM CuO [14]. Surprisingly, we find a non-zero mixed signal not only in the AFM, but also in the PM phase with nearly the same intensity. It is important to note that we are not considering a Curie paramagnet (such as Ni above \( T_C \) with disordered and/or fluctuating magnetic moments, but a Pauli PM state, where the magnetization is strictly zero in all points of space. Therefore, our finding that the mixed signal is essentially unchanged when going from the AFM to the PM state unambiguously proves that it is unrelated to local magnetic moments, in contrast to the interpretation in Refs [14, 16].

Rather than being of magnetic origin, the non-zero mixed signal is in fact induced by angular momentum transfer from the light helicity to the electron spin via SO in the core shell together with a strong exchange
effect in the decay process. To see this, consider light with left (+) helicity and a non-magnetic ground state. The $2p_{3/2}$ optical transition has a larger amplitude for spin-up than for spin down electrons because of the dominantly parallel alignment of spin and orbit in $2p_{3/2}$. For example, for an empty or spherically symmetric 3$d$ shell the intensity ratio is 5:3. Consider now a spin-up electron transition. The RPES intermediate state has one extra spin-up electron in the 3$d$-shell (denoted $u^\uparrow$) and a $2p$-hole of dominant spin-up character. This state decays through Coulomb interaction to the photoemission final state with one 3$d$-hole and the photoelectron. The direct Coulomb matrix elements is of the form $\langle k\sigma, c\uparrow |V| v\sigma, u^\uparrow \rangle$ which is independent of the photoelectron spin $\sigma$. So the direct decay alone would lead to a spin-balanced photocurrent. For the exchange decay, the matrix element is $\langle k\sigma, c\uparrow |V| v\uparrow, v\sigma \rangle \sim \delta(\sigma, \uparrow)$. This is roughly as large as the direct Coulomb term for spin-up electrons (the radial matrix elements are exactly the same) but it is zero for spin-down electrons. Since the exchange matrix elements are subtracted from the direct terms in Eq. (1), the transition probability for spin-up electron emission is strongly reduced by the exchange process. This shows that a core-valence transition of a spin up electron leads, through autoionization, to a strongly spin polarized photocurrent with a majority of spin down electrons. As mentioned before, left circular polarized light promotes dominantly spin-up electrons in the $2p_{3/2}$-$3d$ transition.

Therefore it produces a majority of spin-down photoelectrons. Under the assumption of complete cancellation between direct Coulomb and exchange matrix elements for parallel spins and by neglecting the direct valence photoemission, the ratio of spin-down to spin-up photoelectrons is $5:3$, which corresponds to a spin-polarization (ratio of mixed over total signal) of $-1/4$. In angle integrated RPES at maximum resonance (Fig. 2a, $h\nu = 554.4$ eV) we find a SP of $-0.21$, in good agreement with such model estimation. These values agree also well with the measured spin-polarization in CuO $^{14}$ and Ni $^{16}$, which is 10–40% depending on binding energy. Our findings clarify the physical mechanism inducing the presence of the mixed signal in both phases, and point to a critical re-examination of experimental observations.

Interestingly, we find that, contrary to the previous set up, it is possible to have a net spin polarization signal on the PM phase. This is possible under appropriate geometrical conditions, and even with unpolarized light. Such SP can be of opposite sign and be due to different active mechanisms. In Fig. 3c, normal emission spectra are shown for light incident along $[1\bar{1}0]$, i.e. perpendicular to the spin-quantization axis $s = e_z$ (perpendicular geometry). As before, the dichroic signal is zero, as light incidence ($p$) and electron emission vector ($n$) lie in a mirror plane of the surface (see Fig. 1a). However, the set up (including spin resolution) is chiral, since the three vectors $p$, $n$ and $s$ form a right-handed frame. Thus SO-induced SP cannot be ruled out by symmetry and a small, positive SP (in this case transverse to the scattering plane) is indeed observed in RPES, even for unpolarized light. A similar SP from PM surfaces for unpolarized light was theoretically predicted in direct PES $^{30}$ in a relativistic approach and confirmed by experiments $^{31, 32}$. It was ascribed to broken symmetry due to the off-normal light incidence together with SO in the initial states and phase shift differences. We do not observe this effect in non-resonant PES since the SO coupling in the Cr 3$d$ valence states is very weak and neglected here. However, for RPES, such SP has to be related to the dynamical SP studied in atomic physics, which is known to be related to phase shift differences in the final outgoing waves, and to be generally small $^{33, 34}$. Our result confirms that such SP exists for an atom embedded in a solid and that it survives to the multiple scattering effects.

A SP signal in the PM phase is also present for parallel geometry with off-normal emission (Fig. 3d). In this case, the system composed by the surface, light incidence (along $e_z$) and electron emission vector, is chiral. Therefore a dichroic signal is observed even in non-resonant PES, known as circular dichroism in angular distribution $^{35}$. In RPES, the angular momentum of the photon is partly transferred to the electron spin through the SO coupling in the $2p$ shell, leading to non-zero intensity also for spin resolved and mixed signals. The spin polarization is negative, i.e. photoelectrons are mainly polarized
antiparallel to their emission direction, because of the exchange process in the autoionization decay. This finding suggests a Fano-like effect in resonant processes for off normal emission directions, which could be well studied along the same lines as direct PES on paramagnets [30].

In conclusion, we have presented a first-principles approach for RPES in solids and its application to Cr(110). By comparing Pauli PM and AFM states, we have shown new effects in normal emission directions, which could be well studied across the light helicity. New effects in the SP suggest that a mapping of spin interactions in paramagnets and disordered magnetic structures could be obtained via full tomography experiments at the core resonances even with unpolarized light.

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