Theory of quasiparticle spectra for Fe, Co, and Ni: bulk and surface

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The correlated electronic structure of iron, cobalt and nickel is investigated within the dynamical mean-field theory formalism, using the newly developed full-potential LMTO-based LDA+DMFT code. Detailed analysis of the calculated electron self-energy, density of states and the spectral density are presented for these metals. It has been found that all these elements show strong correlation effects for majority spin electrons, such as strong damping of quasiparticles and formation of a density of states satellite at about − 7 eV below the Fermi level. The LDA+DMFT data for fcc nickel and cobalt (111) surfaces and bcc iron (001) surface is also presented. The electron self-energy is found to depend strongly on the number of nearest neighbors, and it practically reaches the bulk value already in the second layer from the surface. The dependence of correlation effects on the dimensionality of the problem is also discussed.

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Introduction

The late 3d metals iron, cobalt, and nickel and their compounds are vital for nearly all fields of technology. The Earth core is believed to be composed predominantly of iron. It is ironic that in the early 21-st century we still lack a complete understanding of these metals. Properties of “normal” (weakly correlated) solids are described quantitatively by density functional theory (DFT) in the local density (LDA) or generalized gradient (GGA) approximations. Not only DFT provides the ground state properties, but in many cases it also gives a rather good description of excitation spectra in terms of Kohn-Sham quasiparticles. The concept of quasiparticles originates from Landau’s Fermi liquid theory and for weakly correlated solids the quasiparticles (electrons and holes) are well defined in a wide energy range.

Fe, Co and Ni, however, are more correlated systems. They have partially filled shells of fairly localized 3d electrons. These electrons form a narrow d-band, and their behavior shows signs of both atomic-like and free-electron-like behavior. In strongly correlated systems the quasiparticle picture breaks down, except in a close vicinity of the Fermi surface. Quasiparticles often have short lifetimes and therefore are not well defined, and in many cases incoherent features such as Hubbard bands and satellites appear in excitation spectra. LDA and GGA typically fail for this class of systems and their theoretical description remains a great fundamental challenge.

In particular, LDA and GGA give rather good magnetic moments for Fe, Co, and Ni, but fail to describe their electronic structure adequately. In particular, photoemission experiments for these metals demonstrate that LDA/GGA calculations give too wide majority spin 3d band, overestimate the spin splitting and fail to reproduce the 6 eV satellite in nickel, an essentially incoherent feature. Some other theoretical methods are needed to properly describe the electronic structure of Fe, Co and Ni.

One of the successful schemes for correlated electron systems is the Dynamical Mean-Field Theory (DMFT), which replaces the lattice problem with a problem of a single correlated site in a self-consistent bath (impurity problem). It has been originally developed for the Hubbard model. Being \( W \) the bandwidth and \( U \) the Coulomb interaction, the DMFT catches the main features of weakly \( (W \gg U) \), intermediate \( (W \sim U) \) and strongly \( (W \ll U) \) correlated regimes, and becomes exact in the limit of infinite dimensions. The crucial point of the DMFT is in the solution of the self-consistent impurity problem. The choice of the DMFT “solver” for a given system is always a compromise between generality, accuracy and efficiency. There exist both numerically exact solvers (quantum Monte-Carlo, exact diagonalization) which can in principle be applied to all systems, and approximate solvers with limited area of applicability but high efficiency, such as the Spin Polarized T-matrix Fluctuation exchange (SPTF) solver for the case \( W \lesssim U \).

Although DMFT was designed originally for the Hubbard model, it can be combined with LDA to describe realistic materials with a local electron correlation. This approach, known as LDA+DMFT, is at present the most universal practical technique for calculating the electronic structure of strongly correlated solids. LDA+DMFT has been successfully applied to various important problems, including, e.g., the electronic structure of manganese, \( \delta \)-Pu, and the \( \alpha - \gamma \) transition in cerium, and the metal-insulator transition in \( \text{V}_2\text{O}_3 \). Despite all success stories of LDA+DMFT, the method is still less than a decade old and at a stage of active development. Most available implementations apply some drastic simplifications to the LDA+DMFT formalism. In particular, many LDA+DMFT codes are based on atomic sphere approximation (ASA)-based LDA codes (LMTO-ASA or
FIG. 1: (Color online) Imaginary part of the self energy $\text{Im}\Sigma(\epsilon + i0)$ for bcc iron, majority spin for orthogonalized LMTO (ORT) and muffin-tin-only (MT) correlated subspaces. Two different temperatures are considered: $T = 400$ K and $T = 1500$ K.

FIG. 2: (Color online) Imaginary part of the self energy for bcc iron, minority spin for two different correlated subspaces (MT and ORT orbitals) and two different temperatures ($T = 400$ K and $T = 1500$ K).

FIG. 3: Real part of the self energy for bcc iron (MT orbitals, $T = 1500$ K).

**I. LDA+DMFT METHOD**

**A. Correlated subspace**

The LDA+DMFT method defines a "correlated subspace" $\{|\mathbf{R}, \xi\rangle\}$ of the strongly correlated orbitals $|\mathbf{R}, \zeta\rangle$, where $\mathbf{R}$ stands for the Bravais lattice site and the quantum number $\xi$ specifies the correlated orbitals within the unit cell. Within this subspace the many-body problem is solved in a non-perturbative manner using DMFT. All remaining states of the crystal are assumed weakly correlated and treated within LDA. For simplicity, we can always choose correlated orbitals to be orthogonal and normalized $\langle \mathbf{R}_1, \xi_1 | \mathbf{R}_2, \xi_2 \rangle = \delta_{\mathbf{R}_1, \mathbf{R}_2} \delta_{\xi_1, \xi_2}$.

Results of a LDA+DMFT calculation depend on the choice of the correlated orbitals. The correct form of $|\mathbf{R}, \xi\rangle$ is dictated by physical considerations for each par-
particular problem\textsuperscript{23,24,25}. Usually the correlated orbitals are derived from \(d\) or \(f\) atomic states. In this case \(\xi\) stands for site (within unit cell) and atomic quantum numbers \(l, m, \sigma\). In early LDA+DMFT implementations \(|R, \xi\rangle\)’s were taken as orthogonalized muffin tin orbitals (MTO’s). Apart from the technical simplicity, however, there is no reason for making such a choice. In particular, orthogonalized MTO’s are poorly localized in real space (due to orthogonalization) and they don’t have pure \(lm\) character (due to tail cancellation and orthogonalization). Besides, such choice of \(|R, \xi\rangle\) de facto restricts the LDA part of the LDA+DMFT code to muffin-tin based methods (LMTO, NMTO, KKR) with a minimal basis set. Nowadays other choices, such as Wannier-like orbitals\textsuperscript{23,25} are investigated. Apart from atomic states, the correlated orbitals \(|R, \xi\rangle\) can be chosen as hybridized orbitals (possibly describing covalent bonds) if dictated so by the physical problem. Sometimes the LDA Hamiltonian is downfolded in order to include only correlated degrees of freedom (such as \(d\)-states or even \(t_{2g}\) or \(e_g\) states). This results in a very time efficient DMFT implementation, however this approach cannot be used to study the hybridization between sp-electrons and the correlated ones, which is important, e.g., for the superexchange interaction. In this section, we present the LDA+DMFT formalism in a more general form, without making any restrictions on the choice of the correlated subspace \(|R, \xi\rangle\) or the basis set used by the LDA part of the code. We are going to return, however, to the question of choosing the correlated subspace in the section describing our implementation.

In LDA a solid is described by the one-particle Kohn-Sham equation

\[
(H_{LDA} - E)|\psi\rangle = 0, \tag{1}
\]

where the LDA Hamiltonian \(H_{LDA}\) has one-electron form

\[
H_{LDA} = \sum_i h_{LDA}(r_i), \tag{2}
\]

with \(h_{LDA}\) acting in the Hilbert space of one-electron states in a periodic crystal, and the index \(i\) numbering all electrons in the crystal. The LDA+U Hamiltonian adds explicit Coulomb term for the correlated orbitals \(|R, \xi\rangle\) to the LDA Hamiltonian

\[
H_{LDA+U} = H_{LDA} + \frac{1}{2} \sum_{R} \sum_{\xi_1, \xi_2, \xi_3, \xi_4} U_{\xi_1, \xi_2, \xi_3, \xi_4} c_{R, \xi_1} c_{R, \xi_2} c_{R, \xi_3} c_{R, \xi_4}. \tag{3}
\]

It has a form of a multiband Hubbard Hamiltonian with the LDA Hamiltonian used as "hopping". The Coulomb parameters \(U_{\xi_1, \xi_2, \xi_3, \xi_4}\) are screened Coulomb integrals for the states \(|R, \xi\rangle\). They are to be found empirically or calculated from first principles, and they, in general, depend on the choice of the correlated subspace \(|R, \xi\rangle\). Note that \(H_{LDA}\) is not a kinetic energy and the Hubbard-U term is not a "raw" Coulomb interaction. Rather, the LDA+U Hamiltonian is an effective Hubbard Hamiltonian for the correlated orbitals, and the rest of the states (e.g. sp states) is described within LDA. The justification of this approach\textsuperscript{24} is not a trivial matter, and the main practical reason why it is widely used is the local nature of the screened Coulomb interaction. Note that we have included the explicit Coulomb term in the Hamiltonian, although many static effects of the Coulomb interaction are already included in the LDA Hamiltonian. Namely, LDA includes a Hartree term, an exchange term, and some correlation effects (including a good description of the screening). However, the many-body treatment of the Hubbard-U part of the Hamiltonian will also give the Hartree-Fock term and various correlation terms. Therefore, a double-counting correction scheme is necessary

\[
\Sigma(z) \rightarrow \Delta \Sigma(z) \equiv \Sigma(z) - \Sigma_{dc}, \tag{4}
\]

where \(\Sigma(z)\) is the local self energy of the DMFT problem. For treating metals within LDA+DMFT the most common choice of the double-counting correction is the static part of the self-energy:

\[
\Sigma_{dc} = \Sigma(+i0). \tag{5}
\]
The multiband Hubbard Hamiltonian, such as Eq. 3 cannot be solved exactly, therefore various approxima­tions are applied. Unlike the static LDA+U method, LDA+DMFT takes into account dynamical corelation ef­fects through the frequency-dependent self energy $\Sigma(z)$.

**B. LDA+DMFT equations**

Spectral density functional theory uses the local Green function $G_R(z)$ as the main observable quantity, much like the particle density $\rho(r)$ in DFT or the one-electron Green function $G(z)$ in Baym-Kadanoff theory. Namely, for sensible definitions of $G_R(z)$, a functional $\Gamma[G_R]$ exists, which is minimized by the true value of $G_R(z)$, and the local self energy $\Sigma_R(z)$ plays the role similar to the Kohn-Sham potential in DFT. In the present paper we define $G_R(z)$ as the projection of the total one-electron GF to the correlated states $|\mathbf{R}, \xi\rangle$ of a given site $\mathbf{R}$

$$G_R(z) = P_R G(z) P_R,$$

where

$$P_R = \sum_{\xi} |\mathbf{R}, \xi\rangle \langle \mathbf{R}, \xi|$$

is the projection operator to the correlated subspace belonging to site $\mathbf{R}$. The one-electron GF $G(z)$ can in turn be expressed via the one-electron self energy $\Sigma(z)$ as

$$G(z) = \left([z - \mu] - h_{LDA} - \Sigma(z)\right)^{-1},$$

where $\mu$ is the chemical potential, and $h_{LDA}$ plays the role of the unperturbed Hamiltonian (“hopping”).

Precisely like in DFT or Baym-Kadanoff theory, the exact expression for the functional $\Gamma[G_R]$ is not known. The most widely used approximation is the dynamical mean field theory (DMFT). The approximation behind DMFT is that the total one-electron self energy $\Sigma(z)$ is taken as the sum of the local self energies of all lattice sites (with double-counting correction when appropriate)

$$\Sigma(z) = \sum_{\mathbf{R}} \Sigma_R(z).$$

This self energy is local, i.e. it does not have matrix elements between different sites

$$\langle \mathbf{R}_1, \xi_1 | \Sigma | \mathbf{R}_2, \xi_2 \rangle = \delta_{\mathbf{R}_1, \mathbf{R}_2} \langle \Sigma_R \rangle_{\xi_1, \xi_2}.$$

With this form of $\Sigma(z)$, the local self-energy $\Sigma_R(z)$ can be obtained from the impurity problem, with the rest of the lattice replaced by the bath GF (or “dynamical mean field”) $G_0^{-1}(\mathbf{R}, z)$ defined by

$$G_0^{-1}(\mathbf{R}, z) = G_R^{-1}(z) + \Sigma_R(z).$$

In a periodic solid all atoms are equivalent, therefore the local quantities such as $\langle \Sigma_R \rangle_{\xi_1, \xi_2}$, $G_R(\xi_1, \xi_2)$ and $G_0^{-1}(\mathbf{R}, z)$ take a block diagonal form with a block for each site.

The system of one correlated site in the self-consistent bath does not have a Hamiltonian, but can be described by an effective action. Here and in the following we use the Matsubara formalism for a finite temperature $T$. The
The action in the Matsubara formalism is

\[
S = -\int dx_1 dx_2 c^\dagger(x_1) \mathcal{G}_0^{-1}(x_1, x_2) c(x_2) + \frac{1}{2} \int dx_1 dx_2 dx_3 dx_4 c^\dagger(x_1) c^\dagger(x_2) U(x_1, x_2, x_3, x_4) c(x_4) c(x_3),
\]

set that satisfies the Bloch theorem: for any translation vector \( \mathbf{T} \)

\[
\psi_{k, \chi}(r + \mathbf{T}) = e^{i \mathbf{k} \cdot \mathbf{r}} \psi_{k, \chi}(r),
\]

where \( \mathbf{k} \) belongs to the Brillouin zone. There is a one-to-one correspondence between real-space and \( \mathbf{k} \)-space basis sets, given by the Fourier transformation

\[
|k, \chi\rangle = \sum_R e^{i \mathbf{k} \cdot \mathbf{R}} |R, \chi\rangle, |R, \chi\rangle = \sum_k e^{-i \mathbf{k} \cdot \mathbf{R}} |k, \chi\rangle,
\]

This equation is used to determine the LDA+DMFT chemical potential (Fermi energy), which must produce the correct number of electrons. In the following subsection we show how the DMFT equations can be presented in a given LDA basis set.

C. DMFT equations for a given LDA basis set

The Kohn-Sham eigenfunctions belong to the Hilbert space of one-electron states in a solid. The choice of the basis set in this space is dictated by the method (LMTO, LAPW, ...), and we can use a basis either in the real-space or in the reciprocal-space. The real space basis set \( \{ |R, \chi\rangle \} \) is defined by the wavefunction

\[
\psi_{R, \chi}(r) = \psi_\chi(r - R)
\]

which is typically localized in a small area around the lattice site \( R \). The \( \mathbf{k} \)-space basis set \( \{ |\mathbf{k}, \chi\rangle \} \) is a basis
The LDA k-space basis set \( \{|\mathbf{k}, \chi\}\) should be used in order to calculate the local GF (6), since we know the LDA Hamiltonian matrix \( h_{LDA}(\mathbf{k}) \) and the overlap matrix \( S(\mathbf{k}) \) in this basis set. On the other hand, the DMFT impurity problem is formulated for the correlated sub-space with a real-space basis set \( \{|\mathbf{R}, \xi\}\). Transforming back and forth between the local GF’s and the self energies is thus necessary at each DMFT iteration. Using Eqs. (22)–(28), it is easy to show that Eq. (6) becomes

\[
(G_R)_{\xi_1, \xi_2}(z) = \sum_{\mathbf{k}, \chi_1, \chi_2} \langle \xi_1|\mathbf{k}, \chi_1 \rangle [S(\mathbf{k})(z - \mu) - h_{LDA}(\mathbf{k}) - \Sigma(\mathbf{k}, z)]^{-1}_{\chi_1, \chi_2}(\mathbf{k}, \chi_2|\xi_2),
\]

where \( \Sigma(\mathbf{k}, z) \) is the self energy matrix in the LDA basis \( \{|\mathbf{k}, \chi\}\}. This expression would be exact only if the basis set \( \{|\mathbf{k}, \chi\}\) was complete. It would also be exact if the correlated orbitals \( \{|\mathbf{R}, \xi\}\) belonged to the space spanned by the basis functions \( \{|\mathbf{k}, \chi\}\), like for the orthogonalized MTO’s, since it would mean that \( \{|\mathbf{k}, \chi\}\) is complete within the space of interest. In realistic full-potential calculations the completeness of \( \{|\mathbf{k}, \chi\}\) is a reasonable approximation. The local self energy transformed into the LDA basis is in turn given by

\[
\Sigma_{\chi_1, \chi_2}(\mathbf{k}, z) = \sum_{\xi_1, \xi_2} \langle \mathbf{k}, \chi_1|\xi_1 \rangle \Sigma_{\xi_1, \xi_2}(z) \langle \xi_2|\mathbf{k}, \chi_2 \rangle.
\]

The one-particle excitation spectrum of a system is given by the density of states (DOS)

\[
D(\epsilon) = -\frac{1}{\pi} \text{Tr} \left[ \text{Im} G(\epsilon + i0) \right],
\]

and by the spectral density, which is the k-resolved DOS

\[
A(\mathbf{k}, \epsilon) = -\frac{1}{\pi} \sum_{\chi} \langle \mathbf{k}, \chi|\text{Im} G(\epsilon + i0)|\mathbf{k}, \chi \rangle.
\]

The spectral density generalizes the concept of quasiparticle band structure by allowing quasiparticles to decay, thus introducing smearing of bands. In the absence of self energy it reduces to the usual Kohn-Sham band structure

\[
A_{KS}(\mathbf{k}, \epsilon) = \sum_n \delta(\epsilon - \epsilon_n(\mathbf{k})).
\]

As we already mentioned, typical spectral density has coherent (quasiparticles) features and also possibly non-coherent (dispersionless) ones: Hubbard band satellites. Note that DMFT gives Green function at Matsubara frequencies \( i\omega_n \), while DOS and the spectral density are defined via GF at the \( \epsilon + i0 \) contour. The numerical analytical continuation can be done, for example, using the Padé approximation.

II. IMPLEMENTATION

In this paper we introduce the code BRIANNA, a new LDA+DMFT implementation based on the full-potential linear muffin tin orbital (FP-LMTO) code developed in Ref.28. As we already mentioned, FP-LMTO gives an accurate description of solids within LDA, and the full-potential treatment is especially important for open structures and surfaces. On the other hand, FP-LMTO uses a relatively small basis set, which is convenient for calculating Green functions, since it involves inverting a matrix in the LDA basis set for each Matsubara frequency and k-point. The typical basis set is “double-minimal”, i.e. it contains two basis functions per each site and \( l, m, \sigma \), with different “tail energies”. It is still much smaller than the basis set in the plane wave and augmented plane wave based codes.
There are a few other technical issues worth mentioning here. A DMFT solver typically needs about 1000 or more Matsubara frequencies (above the real axis). Instead of performing matrix inversion in Eq. (29) for every frequency (and every \( \mathbf{k} \)-point), it is a standard technique nowadays to use a smaller (usually logarithmic) mesh in Eq. (29) and, after having calculated the local GF, transform it to the Matsubara mesh using cubic splines. Inverse transformation is applied to the self energy in order to plug it into Eq. (29). All calculations of the present paper use 1024 Matsubara frequencies in the DMFT solver, but only 80 points in Eq. (29). The first 16 of them coincide with the first Matsubara frequencies, while the rest forms a logarithmic mesh. Another issue involves numerical analytical continuation in (31) and (32). Analytical continuation using the Pade approximation can introduce serious numerical errors. In the present implementation we do not apply the Pade approximation to the Green functions. Instead, we use it only for the self energy \( \Sigma(z) \), while the GF is calculated directly at the \( \epsilon + i\delta \) contour using Eq. (8).

We have already mentioned the very important question of choosing the orbitals spanning the correlated subspace. In this paper we use two different definitions, both of them atomic-like and derived from transition metal d states. The first, more traditional, uses orthogonalized d-type basis functions of the FP-LMTO method, transformed to the real space via (19). We call this definition orthogonalized LMTO (ORT) correlated subspace. We remind the reader that it is poorly localized, and also that the orbitals \( |\mathbf{R}, \xi\rangle \) do not have pure \( l, m \) character. Such kind of approach requires minimal LDA basis set for the d-type electrons. This FP-LMTO code\(^{28}\) allows to use a double-minimal basis set (two or even more energy tails) for sp-electrons and a minimal one (single tail) for d electrons. The latter have less dispersion than sp electrons, and use of the single-tail basis set for them does not lead to any severe errors within LDA.

Our second choice is somewhat opposite, since it deals with extremely localized correlated orbitals. We call it muffin-tin only (MT) correlated subspace. \( |\mathbf{R}, \xi\rangle \) is chosen as

\[
\Psi_{\mathbf{R},\xi}(\mathbf{r}) = \begin{cases} 
\Phi(t)(|\mathbf{r} - \mathbf{R}_\xi|)Y_{lm}(\mathbf{r} - \mathbf{R}_\xi), & |\mathbf{r} - \mathbf{R}_\xi| < R_{MT,\xi} \\
0, & |\mathbf{r} - \mathbf{R}_\xi| > R_{MT,\xi}
\end{cases}
\] (33)

\( \mathbf{R}_\xi \equiv \mathbf{R} + \mathbf{r}_\xi \) is the site where the orbital \( |\mathbf{R}, \xi\rangle \) is located and \( R_{MT,\xi} \) is the muffin-tin radius for this site. The pure muffin-tin radial function \( \Phi(t) \) is the solution of the radial Schrödinger equation in the spherically averaged Kohn-Sham potential\(^{28}\), inside the muffin-tin only, for a certain energy \( E_v \).

The correlated orbital in Eq. (33) is zero outside a given muffin-tin, and is thus ultimately local. The correlated orbitals have pure angular momentum character, but, at the same time, they are orthogonal by definition (since they do not overlap). Note that the correlated orbitals obviously do not form a complete basis set within the Hilbert space of one-electron wavefunctions (since the interstitial region is not included at all). This is not a problem, since they are only used to define the Hubbard-U term in the Hamiltonian (3), while the "hopping" term is the LDA Hamiltonian defined using the FP-LMTO basis set, which we assume to be sufficiently complete.

### III. RESULTS

#### A. Iron

The LDA+DMFT self energies of bcc iron have been calculated for \( U = 2.3 \) eV, \( J = 0.9 \) eV. We used both muffin-tin only (MT) and the orthogonalized LMTO (ORT) sets of correlated orbitals, and also performed calculations for different temperatures. In Figures 1 and 2 we present the imaginary part of the self energies,
FIG. 9: (Color online) Spectral density of fcc nickel from LDA+DMFT with MT correlated orbitals at $T = 400$ K.

FIG. 10: (Color online) Density of states of hcp cobalt (LDA+DMFT with MT correlated orbitals at $T = 400$ K vs LDA)

Im $\Sigma(\epsilon + i0)$, for majority and minority spins respectively, while in Figure 3 we show the real part of $\Sigma$ (only for the MT basis and for $T = 1500$ K). In quasiparticle language the matrix elements $Re \Sigma(E + i0)_{\xi,\xi}$ describe the shift of the quasiparticle bands, while $-Im \Sigma(\epsilon + i0)_{\xi,\xi}$ has the physical meaning of quasiparticle band smearing $\Gamma$, which is the inverse of the quasiparticle decay time $\tau = 1/T$. The band structure is only well defined if $|Im \Sigma_{\xi,\xi}| \ll W$, where $W$ is the bandwidth. For metals this is always true in the vicinity of the Fermi level, since $Im \Sigma(\epsilon + i0) = 0$. The self energies presented here are averaged over the orbital indices $m$, namely

$$\Sigma(z) = \frac{1}{5} \sum_{m} \Sigma_{mm}(z). \tag{34}$$

The crystal field splitting of $\Sigma$ is rather small and we are not going to discuss it in details. Figure 4 shows the total density of states (DOS) of bcc iron (LDA+DMFT vs LDA), while the spectral density (k-resolved DOS) is presented in Figure 5 for several high-symmetry directions. We remind the reader that the spectral density is the generalization of the band structure with finite quasiparticles lifetime taken into account. Both figures use the MT basis set and $T = 400$ K.

The three curves in Figs. 1 and 2 are qualitatively similar, proving that both MT and ORT correlated orbitals (corresponding to well-localized and poorly localized d-states, respectively) can be used to adequately describe iron within LDA+DMFT. However, the exact amplitude of the peaks in $\Sigma$ is sensitive to the choice of the correlated subspace. We are going to use the muffin-tin only (MT) correlated orbitals for the rest of this paper. Note also that $\Sigma$ is practically temperature-independent for a wide range of temperatures.

The majority spin $Im \Sigma(\epsilon + i0)$ in Fig. 1 has the main peak at $\epsilon \simeq -7$ eV, by reaching the value $-3.4$ eV. This gives rather strong damping of quasiparticles, as we can observe in Fig. 5. There is also a shoulder or small minimum at $\epsilon \simeq -2$ eV. The correlation effects are more pronounced for the majority spin electrons, which is common for late transition metals (see Ref. 22 for an interesting discussion). The LDA+DMFT density of states (Fig. 4) shows the narrowing of the majority-spin d-band compared to the LDA DOS and also a satellite at $\epsilon \simeq -7$ eV. This is the effect of $Re \Sigma(\epsilon + i0)$. The positive region of $Re \Sigma$ for the majority electrons between $-6$ eV and the Fermi level in Fig. 3 leads to the narrowing of the band, while the sharp negative peak at $-8$ eV "draws" the electrons down in energy, leading to the formation of the DOS satellite. Naturally, the smearing of the quasi-
particle bands, given by $\text{Im} \Sigma$, leads to the smearing of the sharp peaks of the LDA DOS. Note that our self energies and DOS differ somewhat from the ones in Ref. 21. In particular, we clearly observe a DOS satellite at $\epsilon \approx -7$ eV, which was not observed in the earlier calculation for $U = 2.3$ eV and $J = 0.9$ eV, but only for much larger values of $U$. The reason, we believe, is that Ref. 21 used a simplified version of the SPTF solver, while in the present paper the full implementation of the SPTF$^{12}$ is used. To the best of our knowledge this is the first calculation that shows the existence of such a satellite, and this could open a new scientific problem, since it has never been reported in any experiment.

B. Cobalt and nickel

The LDA+DMFT self energies for fcc nickel, hcp cobalt and fcc cobalt are presented in Figs. 6 and 7,
and the bcc iron self energy is also shown for comparison. The values of Hubbard parameters were $U = 2.3$ eV, $J = 0.9$ eV for cobalt and $U = 3$ eV, $J = 1$ eV for nickel. Strictly speaking the parameters $U$ and $J$ are somewhat arbitrary (since they apply to a model LDA+U Hamiltonian) and their values depend on the choice of the correlated orbitals. We make a rather traditional choice of $U$ and $J^{21,22}$ in the present paper, however$^{4,12,21,22}$.

The general structure of the self energy is similar for Fe, Co and Ni. The correlation effects for majority spin electrons are strongest for nickel and weakest for iron, at least for the values of $U$ and $J$ used here. The shoulder at $-2$ eV is most pronounced for iron and practically disappears for nickel. The self energy curves for fcc cobalt and hcp cobalt are almost identical. The correlation for minority spin electrons (Fig. 7) are by far strongest in nickel, which has the lowest magnetic moment of the three elements, therefore the difference between majority and minority spin behavior is less profound in nickel compared to iron and cobalt.
FIG. 17: (Color online) Spectral density of fcc cobalt (111) surface modelled by the 5-layer slab (from LDA+DMFT with MT correlated orbitals at $T = 400$ K).

FIG. 18: (Color online) Imaginary part of the self energy for bcc iron (001) surface modelled by the 5-layer slab, majority spin (from LDA+DMFT with MT correlated orbitals at $T = 400$ K). In the legend "1 of 5" indicates the surface atom of the slab, "2 of 5" the sub-surface atom and "3 of 5" the quasi-bulk atom.

FIG. 19: (Color online) Imaginary part of the self energy for bcc iron (001) surface, modelled by the 5-layer slab, minority spin (from LDA+DMFT with MT correlated orbitals at $T = 400$ K). In the legend "1 of 5" indicates the surface atom of the slab, "2 of 5" the sub-surface atom and "3 of 5" the quasi-bulk atom.

Figures 8 and 10 present density of states of fcc nickel and hcp cobalt respectively, while Figures 9 and 11 show the spectral density for these materials. The density of states for nickel is in a good agreement with previous LDA+DMFT calculations. Note that the SPTF solver places the majority-spin satellite at about $-7.5$ eV, while in experiment it is observed at $-6$ eV. The spectral density of fcc nickel is, to the best of our knowledge, presented here for the first time. Since bcc iron, fcc cobalt and fcc nickel have different crystal structure, their band structures naturally look different. However, Fe, Co and Ni all have strong smearing of majority-spin bands at about $-7$ eV dictated by the peak in the self energy (Fig. 6), and show a DOS satellite at about $-7.5$ eV.

The LDA+DMFT values of the spin magnetic moments are substantially equal to the LDA values (e.g., for bcc Fe we have $\mu = 2.23\mu_B$ per atom from the DMFT calculation which should be compared to $\mu = 2.22\mu_B$ per atom from LSDA, and for hcp Co we obtain $\mu = 1.54\mu_B$ per atom from the DMFT calculation which should be...
FIG. 20: (Color online) Spectral density of bcc iron (001) surface modelled by the 5-layer slab (from LDA+DMFT with MT correlated orbitals at $T = 400$ K).

FIG. 21: (Color online) Real part of the self energy for bcc iron (001) surface modelled by the 5-layer slab. Both majority and minority spins for the surface atom (labelled "1 of 5") are reported and compared to the bulk values (from LDA+DMFT with MT correlated orbitals at $T = 400$ K).

compared to $\mu = 1.57\mu_B$ from LSDA). Indeed the problem of the effect of the correlations on the spin and orbital magnetic moments is very interesting and will be the subject of further investigations in the near future.

C. Surfaces

We model the nickel (111) surface with slabs having different number of close-packed atomic layers. Such slabs form a superlattice with a 30 Å thick layer of vacuum separating them, with each slab having two (111) surfaces. The calculations have been done for 5-layer slabs; however just for methodical aims, to show the sensitivity to the computational details the results for 3-layer slabs are also presented on some figures. In Figs. 12 and 13 we present the LDA+DMFT self energies (imaginary part) for nickel slabs for majority and minority spin respectively. Data for each layer of the 3-layer and 5-layer slabs and for the bulk fcc nickel (for comparison) are presented. Notice that the self energy of a nickel atom at the surface is obviously quite different from the self energies for the rest of the atoms in the slab. The most noticeable effect is that the positions of the peaks are shifted and that the correlation effects for majority spin electrons seem to be more enhanced at the surface compared to bulk. We will encounter similar effects for the other surfaces studied here (see below). This shows that the effect of the correlations is different for the topmost surface layer, compared to the rest of the surface layers, and that the sub-surface layer already seems really bulk-like. This finding is an observation that is worthy experimental attention. The reasons of such a difference are rather obvious: due to the reduced coordination number of the surface atoms the bands become narrower, which makes correlation effects more important. In addition the screening of the electron-electron interaction is less effective for the surface atoms, and this increases the value of the Hubbard U. Although the basic mechanisms are easily identified for why correlation effects are more important at the surface, we provide here a quantitative measure of this effect.

The spectral density of the nickel 5-layer slab is presented in Fig. 14 along high-symmetry directions of the two-dimensional Brillouin zone. For well-defined quasi-
particles, each band of the bulk nickel splits into five bands for the 5-layer slab. Some of the bands are surface states, while the rest joins into the bulk continuum when the number of atomic layers go to infinity. In Fig. 14, it is already possible to observe the surface states (isolated bands) and the hint of the bulk continuum formation (several bands that are very close to each other).

Similar results are obtained for the fcc cobalt (111) surface and for bcc iron (001) surface, whose self energies are respectively shown in Figs. 15-16 and in Figs. 18-19. We can notice two main differences with respect to the results for the nickel: for the majority spin the shift of the peak and the increase of its depth for the atoms on the surface are stronger, while for the minority spin the correlation effects are, somewhat surprisingly, decreased (slightly for Co and strongly for Fe). In Fig. 21 we can observe the real part of the self energies for the atoms on the iron surface, compared to the bulk values. It is espe-
cially interesting that for the surface layer the self energy of minority spin states is considerably suppressed, which leads to the fact that the satellite at $-7.5$ eV is almost totally polarized and possesses majority spin character. In Figs. 17 and 20 we show the spectral densities for the surfaces of Co and Fe, and the effects of the change of the imaginary parts of $\Sigma$ with respect to the bulk are clearly evident in the difference of the definition of the quasi-particle bands for majority and minority spins.

Finally we have to notice that the choice of the values of $U$, already non trivial for the bulk materials, becomes more problematic for the surfaces, where the screening is much smaller. To analyze this problem, we have tried to model the bcc Fe (001) surface with different values of $U$ for atoms belonging to different layers, namely $U = 2.3$ for the inner layer, $U = 2.4$ for the intermediate one and $U = 3.0$ for the external one. In Figs. 23 and 24 we respectively show the real and the imaginary part of the self energy for the external atoms. In comparison to the previous calculation we do not observe any drastic effect, but mainly a reasonable increasing of the peaks and a small shift of the $-7.5$ eV satellite. This makes the satellite more pronounced in the density of states.

In Figs. 17 and 20 we show both bulk and surface states. The question “How do the correlation effects depend on the dimensionality of the problem?” still needs further investigation, however, and the the LDA+DMFT studies of slabs of different thickness and nanowires are the subject of the future research.

IV. CONCLUSION

In this paper we have introduced the new full-potential LDA+DMFT code BRIANNA and we have applied it to the correlated electronic structure of bulk Fe, Co, Ni, and the fcc Co and Ni (111) surface, and the bcc Fe (001) surface. The calculated self energies, DOS and the spectral densities are presented. The spectral density plots show the correlated electronic structure in the most clear way, as the $k$-resolved DOS (or, equivalently, as the smeared band structure). The main correlation effects in iron, cobalt and nickel are observed for the majority spin electrons and they include strong quasiparticle damping for at about $-7$ eV, narrowing of the d-band (compared to LDA/GGA) and the appearance of a DOS satellite at about $-7.5$ eV, which is a non-quasiparticle feature.

The calculations for Ni and Co (111) surfaces and for Fe (001) surface show that the electron self energy depends mostly on the local coordination number, with the atoms in the second layer from the surface already being similar to the bulk. Hence our calculations suggest that the effect of correlations should be different for the surfaces of these elements, compared to the bulk. In addition, the spectral density of the Ni (111) surface show both bulk and surface states. The question “How do the correlation effects depend on the dimensionality of the problem?” still needs further investigation, however, and the the LDA+DMFT studies of slabs of different thickness and nanowires are the subject of the future research.

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19 K. Held, G. Keller, V. Eyert, D. Vollhardt, and V. I. Anisi-