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#### Co(CO)n/Cu(001)

Towards understanding chemical control of the Kondo effect Bahlke, Marc Philipp; Wahl, Peter; Diekhöner, Lars; Herrmann, Carmen

Published in: Journal of Applied Physics

DOI (link to publication from Publisher): 10.1063/1.5079518

Publication date: 2019

**Document Version** Publisher's PDF, also known as Version of record

Link to publication from Aalborg University

Citation for published version (APA): Bahlke, M. P., Wahl, P., Diekhöner, L., & Herrmann, C. (2019). Co(CO)n/Cu(001): Towards understanding chemical control of the Kondo effect. *Journal of Applied Physics*, *125*(14), Article 142910. https://doi.org/10.1063/1.5079518

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# Co(CO)<sub>n</sub>/Cu(001): Towards understanding chemical control of the Kondo effect

Cite as: J. Appl. Phys. **125**, 142910 (2019); https://doi.org/10.1063/1.5079518 Submitted: 31 October 2018 . Accepted: 14 February 2019 . Published Online: 04 April 2019

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Cite as: J. Appl. Phys. **125**, 142910 (2019); doi: 10.1063/1.5079518 Submitted: 31 October 2018 · Accepted: 14 February 2019 · Published Online: 4 April 2019



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#### ABSTRACT

The Kondo effect is a many-body phenomenon, allowing insight into the electronic and atomistic structure of magnetic adsorbates on metal surfaces. Its chemical control is intriguing because it deepens such insight, but the underlying mechanisms are only partly understood. We study the effect of increasing the number of CO ligands attached to a cobalt adatom on copper(001), which correlates with an increase in the Kondo temperature  $T_K$  experimentally [Wahl *et al.*, Phys. Rev. Lett. **95**, 166601 (2005)], by solving an Anderson impurity model parametrized by the density functional theory. Our results suggest that the orbital responsible for the Kondo effect is  $d_{x^2-y^2}$  for the tetracarbonyl and its combination with  $d_{z^2}$  for the dicarbonyl. The molecular structures depend considerably on the approximate exchange–correlation functional, which may be related to the known difficulty of describing CO binding to metal surfaces. These structural variations strongly affect the Kondo properties, which is not only a concern for predictive studies but also of interest for detecting mechanism ing the tetracarbonyl to  $C_{4\nu}$  symmetry, as suggested by experimental data, we find structures compatible with the experimental trend for  $T_K$  (employing BLYP-D3+U). This is not possible for the tricarbonyl despite the range of computational parameters scanned. For the tetra- and dicarbonyl, the increased  $T_K$  correlates with a larger hybridization function at the Fermi level, which we trace back to an increased interaction of the Co 3*d* orbitals with the ligands.

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#### I. INTRODUCTION

When a magnetic atom or molecule is brought into contact with a metallic system, the conduction-band electrons can screen the spin of this atom or molecule, due to strong electron correlation. If such screening takes place in impure metallic systems, the resistivity starts to grow with decreasing temperature, as opposed to its pure counterpart, which has been first explained by Kondo for dilute magnetic alloys in  $1964^{1-3}$  (the name impurity has later been extended to include magnetic adsorbates as well). Since then, the Kondo effect has become a subject of intensive research since it allows insight into fascinating aspects of electron correlation<sup>4</sup> and can give information on the electronic and atomistic structure of molecular adsorbates.<sup>5-14</sup>

These strong correlations lead to collective low-energy excitations close to the Fermi surface. Below a certain temperature, impurity electrons dressed by these excitations can behave as weakly interacting quasiparticles in the spirit of Landau's phenomenological Fermi-liquid theory. It was Nozières who showed that the Kondo effect can be characterized in terms of a so-called local Fermi liquid in 1974, which was a big step towards conceptual understanding of Kondo systems.<sup>15</sup>

Magnetic molecules on metallic substrates are of particular interest for spintronics, because of their potential to self-assemble on a surface.<sup>5,9,16</sup> The magnetic character of the molecules can be used for storing data or also for transferring information via spin coupling between different neighboring spin centers.<sup>17</sup> For applications in spintronic devices, an understanding is required of the coupling of the adsorbate's magnetic moment not only with neighboring spin centers but also with conduction-band electrons provided by the substrate. As mentioned earlier, the latter can give rise to the Kondo effect and can be controlled by changing the environment of a spin center. For instance, self-assembly of O<sub>2</sub> molecules

leads to periodically enhanced Kondo resonances,  $^{16}$  and by changing the chain length of  $\rm Mn_xFe$  on a  $\rm CuN_2$  surface, the Kondo coupling can be controlled.  $^{13}$ 

In this work, we would like to focus on the effect of molecular ligands on the Kondo effect.<sup>18-21</sup> Wahl et al.<sup>22</sup> have shown that the coupling of the local moment on a Co atom to the conduction-band electrons of a Cu(001) substrate can be enhanced by attaching CO ligands to the Co atom [forming  $Co(CO)_n$  complexes], by extracting the Kondo temperature from scanning tunneling spectroscopy (STS) experiments. The number of CO ligands might not only affect the hybridization of the Co atom but will also change the splitting of the 3d shell, as a consequence of the different symmetries of the complexes. For n = 2, a fourfold symmetry was found in scanning tunneling microscopy (STM), which is due to a thermally induced rotation of the intrinsically twofold symmetric adsorbate on Cu(001), which happens on a faster time scale than what the STM can resolve. Co(CO)<sub>3</sub>/Cu(001) is observed to have no rotational  $(C_{3\nu})$  symmetry, whereas the complex with four ligands exhibits  $C_{4\nu}$  symmetry (which could result both from an intrinsically  $C_{4\nu}$ -symmetric structure and from a rotating  $C_{2\nu}$ -symmetric one, with two opposing ligands being closer to the surface than the other two). The Kondo temperatures  $T_{\rm K}$ , as extracted from a Fano fit of the STS spectra, increase with the number of CO ligands: 165  $\pm$ 21 K (n = 2), 170  $\pm$  16 K (n = 3), and 283  $\pm$  36 K (n = 4).

We want to gain insight into this behavior as a step towards establishing structure-property relationships for the Kondo effect, by solving the Anderson impurity model parametrized by the Kohn-Sham density functional theory (so-called DFT++ methods, following Ref. 23). In doing so, we will point out how shortcomings in present-day first-principles electronic-structure methods when predicting the atomistic structures of adsorbates on surfaces can strongly affect predicted Kondo properties. This is particularly relevant for systems with CO ligands, since interaction of CO with metal surfaces poses a challenge to electronicstructure methods ("CO-puzzle").<sup>24</sup>

#### II. METHODOLOGY

The optimization of molecular structures on surfaces, in the scope of the Kohn-Sham DFT (hereinafter referred to as DFT for the sake of brevity), requires a choice of the approximate exchange-correlation functional that can describe the electronic structure not only of the molecule/adsorbate but also of the metallic substrate to predict adsorption distances, angles, adsorption sites, and symmetries accurately. For molecular adsorbates with their large number of atoms in the unit cell, one often chooses local density approximation (LDA) and generalized-gradient approximation (GGA) type exchange-correlation functionals, as they are a good compromise between accuracy and computational effort in practice. These classes of exchange-correlation functionals are problematic for the description of the CO ligands, due to an underestimated gap between the highest occupied and the lowest unoccupied molecular orbital (HOMO and LUMO). This contributes to the well-known problem of DFT in predicting the correct adsorption sites of CO molecules on different metal substrates, which is known as the "CO-puzzle" in the literature.

The work of Alaei *et al.*<sup>28</sup> shows that using BLYP (a GGA functional)<sup>29</sup> can at least solve the problem of predicting the

correct adsorption sites for CO on some metal substrates, such as Rh(111), Pt(111), and Cu(111), but without solving the problem of the underestimated HOMO-LUMO gap of a CO molecule. On the other hand, the work of Favot *et al.*<sup>30</sup> shows that PBE<sup>31,32</sup> (a GGA functional) is able to predict the correct adsorption site of CO on a Cu(001) surface. A more systematic improvement can be reached by taking into account nonlocal correlation effects,<sup>33</sup> as in the scope of the van der Waals-density functional (vdW-DF with revPBE) developed by Dion *et al.*,<sup>34</sup> although its generalizations to spin-polarized systems<sup>35,36</sup> are not broadly available in electronic-structure codes. Therefore, we focus on spin-polarized PBE and BLYP here for structure optimizations, including their DFT+*U* variants as we have found this to mimic the effect of strong correlation on adsorption distances.<sup>37</sup> Furthermore, we applied Grimme's dispersion correction (DFT-D3) in all cases.<sup>38,39</sup>

All Kohn-Sham DFT calculations were performed with the Vienna ab initio simulation program (VASP) using the projector augmented-wave method.<sup>40-42</sup> For the carbonyl cobalt complexes under study here, we modeled the Cu(001) surface with a super cell size of  $4 \times 4$  Cu atoms with five Cu layers in total. Structural relaxation was done with a  $2 \times 2 \times 1$  k-grid and convergence criteria for the self-consistent field algorithm of  $2.7 \cdot 10^{-5}$  eV, and 0.027 eV/Å for the force acting on each atom. The kinetic-energy cutoff for the plane-wave single-particle basis was set to 400 eV. In addition to the adsorbates, we allowed the two topmost Cu layers to be relaxed and also the cell shape (i.e., the lattice parameter). The relaxed lattice parameter is 3.48-3.49 Å for all systems under investigation, consistently for all optimization protocols. In addition, we optimized the systems (adsorbate and the two topmost Cu layers) with a fixed lattice parameter of 3.615 Å,43 which is denoted by the index "fix." For DFT+U, we used an on-site Coulomb potential of U = 4.0 eV and J = 0.9 eV applied on the Co 3*d* orbitals.

The electronic structure of the optimized carbonyl cobalt complexes on Cu(001) was then analyzed with a combination of the density functional theory and the Anderson impurity model (AIM). For this purpose, we calculated the electronic structure with the spin-unpolarized DFT (as usual in the literature) using the PBE exchange–correlation functional and a *k*-grid of  $17 \times 17 \times 1$  centered around the  $\Gamma$ -point. A slightly more detailed description of the methodology is given in Ref. 37. The Anderson Hamiltonian reads

$$\begin{split} \hat{H} &= \sum_{\nu\sigma} \boldsymbol{\epsilon}_{\nu} \hat{c}^{\dagger}_{\nu,\sigma} \hat{c}_{\nu,\sigma} + \sum_{\nu i\sigma} \left[ V_{\nu i} \hat{c}^{\dagger}_{\nu,\sigma} \hat{d}_{i,\sigma} + V^{*}_{\nu i} \hat{d}^{\dagger}_{i,\sigma} \hat{c}_{\nu,\sigma} \right] \\ &+ \sum_{i\sigma} \boldsymbol{\epsilon}_{i} \hat{d}^{\dagger}_{i\sigma} \hat{d}_{i\sigma} + \frac{1}{2} \sum_{\substack{ijkl\\\sigma\sigma'}} U_{ijkl} \hat{d}^{\dagger}_{i\sigma} \hat{d}^{\dagger}_{j\sigma'} \hat{d}_{l\sigma'} \hat{d}_{k\sigma}, \end{split}$$
(1)

where  $\epsilon_i$  is the energy of the *i*th localized *d* orbital of the impurity (here defined as the Co atom) and  $\epsilon_v$  is the kinetic energy of the bath electron *v* (in this work, the remainder of the system, i.e., the CO ligands and the copper surface).  $\hat{c}_{v\sigma}/\hat{c}_{v\sigma}^{\dagger}$  are creation and annihilation operators for electrons with spin  $\sigma$  acting on the *v*th bath state, whereas  $\hat{d}_{i\sigma}/\hat{d}_{i\sigma}^{\dagger}$  are the corresponding operators acting on the local orbital *i*. The bath electrons are coupled to the impurity via the hybridization  $V_{vi}$ , and  $U_{ijkl} = \int dr dr' \psi_i^*(r) \psi_j^*(r') \frac{e^2}{|r-r|} \psi_k(r) \psi_l(r')$  is the local Coulomb interaction (we dropped the spin indices here for simplicity) as introduced by Slater,<sup>44</sup> with  $\psi_x$  (x = i, j, k, l) being, in general, any atom-centered basis function. We used the parameters  $(F^0, F^2, \text{ and } F^4, \text{ see below})$  as derived by Slater for hydrogen-type atomic orbitals.

The central quantity extracted from a Kohn–Sham DFT wavefunction is the hybridization  $V_{vi}$ , which we employ to compute an energy-dependent hybridization function  $\Delta_{ii}(\omega)$ ,

$$\Delta_{ij}(\omega) = \sum_{\nu} \frac{V_{\nu i} V_{\nu j}^*}{\omega + i0^+ - \epsilon_{\nu}}.$$
(2)

In practice, this was done by projecting the Kohn–Sham Green's function  $G_{\nu\nu'}^{KS}$  onto a set of localized atomic orbitals and then extracting  $\Delta_{ij}(\omega)$  from the local noninteracting Green's function  $g_{ij}(\omega)^{45}$  as

$$\Delta_{ij}(\omega) = -[g_{ij}^{-1}(\omega) + \boldsymbol{\epsilon}_{ij} - (\omega + 0^+)\delta_{ij}]. \tag{3}$$

Here,  $\epsilon_{ij}$  are the matrix elements of the Kohn–Sham operator in the local basis that has been used in the projection. Diagonalization of the local Kohn–Sham operator leads to the impurity levels  $\epsilon_i = \epsilon_{ii}$  introduced in Eq. (1), and consequently  $g_{ij}(\omega) = g_i(\omega)$ , as well as  $\Delta_{ii}(\omega) = \Delta_i(\omega)$ .

The energy-dependent hybridization function was then used to solve the Anderson impurity model with the continuous-time quantum Monte Carlo (CT-QMC) impurity solver in the hybridization expansion<sup>46,47</sup> as implemented in IQIST.<sup>48</sup> The imaginary part of  $\Delta_{ij}(\omega)$  carries the information on the impurity level being broadened by the interaction with the bath electrons, while the energy shift caused by this interaction is captured in the real part of  $\Delta_{ij}(\omega)$ . The term "noninteracting" in this context refers to the DFT solution (although electron correlation effects are captured in the exchangecorrelation term), because it is used to parameterize the AIM. The corresponding "interacting" solution is subsequently obtained from DFT++, which yields the interacting impurity Green's function  $G^{IMP}$ that is connected to the noninteracting one via the Dyson equation,

$$g_{ij}^{-1}(\omega) = G_{ij}^{\text{IMP}^{-1}}(\omega) + \Sigma_{ij}(\omega).$$
(4)

Here,  $\Sigma$  is the so-called self-energy, which describes the dressing of impurity electrons by low-energy excitations caused by electron correlation. This leads to a quasiparticle description of the impurity electrons,<sup>49</sup> which may below a certain temperature behave as a Fermi liquid.<sup>15,50</sup> At this point, it is worthwhile to mention that the self-energy as obtained from our DFT++ approach captures the electron correlation effects on the impurity orbitals explicitly.<sup>51</sup> For this reason, DFT++ works well for strongly correlated materials regardless of the correlation strength, in contrast to present-day approximate DFT, making DFT++ suitable for describing Fermi-liquid-like systems.

The impurity occupation in the interacting case can be obtained by the integration (or rather the Matsubara summation) of the interacting Green's function. The imaginary part is related to the spectral function  $A_{ij}$  as

$$A_{ij}(\omega) = -\frac{1}{\pi} \operatorname{Im} G_{ij}^{\mathrm{IMP}}(\omega), \qquad (5)$$

which requires to be analytically continued<sup>52</sup> onto the real energy

axis (in the simplest case, replacing  $i\omega_n \rightarrow \omega + i\delta$  is sufficient,<sup>53</sup> but due to statistical noise present in CT-QMC, we have to use the maximum entropy method<sup>54</sup>) because CT-QMC operates in imaginary time.

The Coulomb term of Eq. (1) was approximated by using only density-density terms (see Ref. 37 for more details), parameterized by the Slater integrals<sup>44,55</sup>  $F^0$ ,  $F^2$ , and  $F^4$  by using the average Coulomb interaction parameter  $U(F^0) = 4.0 \text{ eV}$  and the exchange-interaction parameter J = 0.9 eV [ $J = \frac{1}{14}(F^2 + F^4) = 0.9 \text{ eV}$  with  $\frac{F^4}{F^2} = 0.625$ ].

The correlation energy already captured in the framework of DFT, often called the double-counting (DC) energy, was corrected by subtracting a term based in the fully localized limit (FLL),<sup>56</sup>

$$E_{\rm DC}^{\rm FLL} = \frac{1}{2} UN(N-1) - \frac{1}{2} JN_{\uparrow}(N_{\uparrow}-1) - \frac{1}{2} JN_{\downarrow}(N_{\downarrow}-1).$$
(6)

In Eq. (6), N,  $N_{\uparrow}$ , and  $N_{\downarrow}$  are the total number of electrons, the number of spin-up, and the number of spin-down electrons on the local subspace, respectively, in our case, the Co 3*d* shell. It is important to note that the fully localized limit is just an approximation to the real correlation energy already captured in the framework of DFT. Thus, in this work, the DC value was shifted by  $\pm 1.0 \text{ eV}$  to evaluate the effect of small errors within the FLL approximation.

Calculating expectation values of operators at finite temperatures typically requires knowledge of the partition function Z of the system.<sup>57</sup> Here, this is achieved by sampling the partition function configurations of the system directly using CT-QMC and employing them in the calculation of observables like  $G^{\text{IMP}}$  or the spinspin correlation function  $\chi$ .<sup>47</sup> The latter is defined on the imaginary time axis (where  $\tau$  runs from 0 to  $\beta$ , with  $\beta = 1/T$ ) as

$$\chi_i(\tau) = \langle \chi_i(\tau)\chi_i(0) \rangle = \langle \hat{S}_Z^i(\tau)\hat{S}_Z^i(0) \rangle.$$
(7)

In Eq. (7),  $\hat{S}_{Z}^{i}(\tau)$  is the local spin at imaginary time  $\tau$ . Evaluating  $\chi(\tau)$ , one can gain insight into the magnetic behavior of the system under consideration. The local spin at the initial time  $\tau = 0$  is equivalent to the magnetization before interactions with the surrounding take place. Another special value which is known as the long-time correlation value of  $\chi(\tau)$  is  $\tau = \beta/2$ , from which one can estimate to which extent the impurity electrons are localized on one of the impurity orbitals, or delocalized/screened due to interaction with the substrate and the ligands. The latter case is indicated by a rather rapid drop of  $\chi(\tau)$  as  $\tau \to \beta/2$ , whereas a finite value at  $\chi(\beta/2)$  suggests that there is a spin density located on one of the impurity orbitals even in the presence of the bath.

#### III. ATOMISTIC STRUCTURE OF Co(CO)<sub>n</sub> ON Cu(001)

Detailed information on bond lengths, bond angles, and adsorption distances is difficult to access from the experiment. At the same time, as discussed above, metal surfaces with molecular adsorbates in general and with adsorbed CO in particular are challenging for present-day first-principles methods. To address this problem, we have optimized all carbonyl cobalt complexes with PBE-D3 and BLYP-D3, with and without U correction. In some cases, we have compared the results of keeping the lattice parameters fixed (for this, we use the index "fix") with the results obtained by relaxing the cell shape together with the atomic positions of the carbonyl complexes (and the surface atoms of the first two layers, see Sec. II for further details). In the following, we briefly summarize the most important observations about the structures of the carbonyl cobalt complexes on Cu(001), while detailed information about the structural parameters can be found in the supplementary material.

#### A. Co(CO)<sub>2</sub>/(001)

For Co(CO)<sub>2</sub>/Cu(001), one consistently obtains  $C_{2\nu}$  symmetry [Fig. 1(a)] for all exchange–correlation functionals mentioned above, which is in agreement with Ref. 22. It can also be confirmed that the rotational barrier of Co(CO)<sub>2</sub>/Cu(001) is rather low (6.4 kJ/mol = 66 meV as obtained from PBE<sub>fix</sub>-D3), which would support the observation of Wahl *et al.*<sup>22</sup> that this molecule is rotating faster than the STM time resolution on Cu(001) in the experimentally accessible temperature range.

Given that present-day approximate DFT has problems to describe the Kondo screening of the magnetic moment, we would expect to obtain a nonzero magnetic moment from spin-polarized DFT optimizations for systems showing a Kondo effect. However, we do not find such a magnetic moment on the adsorbate when using PBE-D3 and BLYP-D3, both for the optimizations including and excluding the cell shape. When we apply a Hubbard U correction on the Co atom for BLYP-D3, we do obtain a magnetic moment of  $1.0\mu_{\rm B}$  on the Co atom of  $\rm Co(CO)_2/Cu(001)$  when we include the cell shape in the optimization. This increased spin localization is in line with the tendency of DFT+U to correct for electron overdelocalization.<sup>58</sup> As shown in Table I, the BLYP-D3+U spin localization has consequences for the adsorption distance as indicated by the increased Co-surface distance  $d_{\text{Co-surf}}$  compared with the BLYP-D3-optimized structure. A similar increase of the adsorption distance can be observed as a result of cell shape relaxation, as suggested by comparing the values for BLYP-D3<sub>fix</sub> (1.39 Å) and BLYP-D3 (1.60 Å). This might come from the slightly reduced lattice parameter (see Sec. II) in cases with cell shape optimization, which effectively decreases the size of the fourfold hollow position and thus increases the adsorbate-surface distance. We will later show that there is a delicate dependence on the Kondo properties obtained from DFT++ on the adsorption distance.

#### B. Co(CO)3/Cu(001)

The DFT-predicted symmetry of  $Co(CO)_3/Cu(001)$  in this study differs from the experimental one reported in Ref. 22 for all exchange–correlation functionals under study. Here, we obtain a rather  $C_{3v}$ -like<sup>59</sup> symmetry of the molecule [see Fig. 1(b)], whereas in Ref. 22, the experimental structure suggests no such rotational symmetry. Of course, one may debate whether the STM results (Fig. 1 in Ref. 22) clearly exclude  $C_{3v}$  symmetry. However, since our optimized structures do not appear consistent with these data, and since we could not find Kondo features in the DFT++ data for any of these structures, we assume that the range of DFT variants employed here is not capable of describing the tricarbonyl structure



**FIG. 1.** (a)  $Co(CO)_2$ , (b)  $Co(CO)_3$ , (c)  $Co(CO)_4$  on Cu(100) ( $C_{4\nu}$ ) and (d)  $Co(CO)_4$  on Cu(100) ( $C_{2\nu}$ ) as obtained from BLYP-D3 + U. We labeled the carbon atoms for  $Co(CO)_3$  and  $Co(CO)_4$  on Cu(100) ( $C_{2\nu}$ ) to point out the two symmetrically different CO ligands.

reliably and will not focus on it any further within the scope of this work (see the supplementary material for more details).

#### C. Co(CO)<sub>4</sub>/Cu(001)

In the case of Co(CO)<sub>4</sub>/Cu(001), it is challenging to obtain a fourfold ( $C_{4\nu}$ )-symmetric structure as suggested by the STM experiments of Wahl *et al.*,<sup>22</sup> because in all attempts to optimize the structure, the twofold ( $C_{2\nu}$ ) symmetry was found to be lower in energy by about 30.2 kJ/mol (PBE<sub>fix</sub>-D3) to 42.1 kJ/mol (BLYP-D3+U).

	PBE-D3 <sub>fix</sub>	PBE-D3	BLYP-D3 <sub>fix</sub>	BLYP-D3	BLYP-D3+U
Co (isolated)	1.46				1.78
Co(CO) <sub>2</sub>	1.33	1.54	1.39	1.60	1.74
Co(CO) <sub>3</sub>	1.58			1.75	1.76
$Co(CO)_4$ ( $C_{2\nu}$ )	1.90			2.07	2.26
$Co(CO)_4 (C_{4\nu})$	1.69			1.89	1.85

TABLE I. Optimized adsorption distances of the carbonyl cobalt complexes on Cu(001) in angstrom as obtained from different DFT protocols. The index "fix" is used in cases where the experimental lattice constant was used.

The vdw-DF functional (revPBE<sub>fix</sub>) predicts  $C_{2\nu}$  symmetry to be 70.0 kJ/mol lower in energy than  $C_{4\nu}$  symmetry. The rotational barrier of the molecule (as obtained for the PBE<sub>fix</sub>-D3 structure) in  $C_{2\nu}$  symmetry is 20 kJ/mol (0.21 eV). This suggests that a hypothetical rotation of this molecule in the STM experiments by Wahl et al. is not responsible for the observed  $C_{4\nu}$  symmetry, because the experiments were taken at T = 6 K (i.e., the thermal energy is not enough to activate a rotation with an energy barrier of 0.21 eV). In the experiment, the preparation of the carbonyl complexes was done by first depositing cobalt on Cu(001), and then saturating the surface with CO molecules. DFT (BLYP+U) suggests a rather short adsorption distance (Table I, also compare Ref. 60) of an isolated Co on Cu(001) (1.78 Å), which is closer to the tetracarbonyl in  $C_{4\nu}$ symmetry (1.85 Å) than to the one in  $C_{2\nu}$  symmetry (2.26 Å). It is conceivable that the formation of the  $C_{4\nu}$ -symmetric system is kinetically favored, due to the adsorption distance being closer to that of an isolated Co on Cu(001). Although the sample was annealed to T = 200 - 300 K, the formation of the probably more stable  $C_{2\nu}$ structure (as suggested by DFT) may be inhibited.

In the DFT and DFT+U calculations, we again do not obtain a local magnetic moment on Co(CO)<sub>4</sub>/Cu(001) in both symmetries. This could make an interpretation in terms of a Kondo effect difficult, but as we will see later, it is indeed possible to identify features in the DFT++ electronic structure for Co(CO)<sub>4</sub>/Cu(001) in  $C_{4\nu}$  symmetry which are in agreement with the experimentally observed Kondo effect. For the corresponding  $C_{2\nu}$  symmetry, no Kondo properties could be found, suggesting that only the  $C_{4\nu}$  symmetry is consistent with the experimental observations.

#### IV. KONDO PROPERTIES OF Co(CO)<sub>n</sub>/CU(001)

In the following, we will focus on the structures optimized with BLYP-D3+U (with optimization of the cell shape), since these show a nonzero local moment on the Co atom for the dicarbonyl complex. For Co(CO)<sub>4</sub> on Cu(001), we limit the discussion to the  $C_{4\nu}$  structure because it fits to the experimentally observed symmetry (see Fig. 1 of Ref. 22) and allows for an interpretation in terms of a Kondo effect (see below). For these structures, we parametrize the AIM with closed-shell PBE, in order to correctly describe the nonmagnetic character of the Cu(001) surface (and thus the coupling of the Co 3*d* orbitals with a nonmagnetic metal).

As the Kondo effect manifests itself as a sharp feature in the spectral function at the Fermi energy, we aim at identifying the relevant Co 3d orbitals that might contribute to the experimentally observed zero-bias anomaly from the spectral functions of the individual orbitals.<sup>22</sup> Nozières<sup>15</sup> showed that a Kondo effect can be

described within the Fermi-liquid theory. We will use this to identify the transition point to the Fermi-liquid regime, as an approximated value for the Kondo temperature  $T_{\rm K}$ . Therefore, we will analyze the temperature dependence of the spin–spin correlation function  $\chi(\tau)$ at the special value  $\tau = \beta/2$ , which should behave as  $T^2$  in the Fermi-liquid regime. Furthermore, we will investigate the so-called first Matsubara-frequency rule,<sup>61</sup> which states that Im $\Sigma(\omega_0)$  should go linearly to zero as  $T \rightarrow 0$  K, as another tool for probing the Fermi-liquid properties of the di- and tetracarbonyl systems.

#### A. Spectral properties of Co(CO)<sub>n</sub>/Cu(001)

In Fig. 2, all Co 3*d* spectral functions of Co(CO)<sub>2</sub> and Co(CO)<sub>4</sub> on Cu(100) ( $C_{4\nu}$ ) are shown at T = 116 K. Due to diagonalization of the Co 3*d* subspace, some of the Co 3*d* orbitals are mixed. Co(CO)<sub>2</sub>/Cu(001) [Fig. 2(a)] shows a sharp feature at the Fermi energy ( $\omega = 0.0$  eV) for the Co  $3d_{0.8z^2 + 0.6(x^2 - y^2)}$  orbital, whereas the remaining Co 3*d* orbitals only have broader features below the Fermi energy (in the energy range shown here). The sharp feature of this orbital thus makes it a promising candidate for causing the experimentally observed Kondo effect (which will



**FIG. 2.** Spectral functions [Eq. (5)] of the Co 3*d* orbitals as obtained from PBE++ for (a) Co(CO)<sub>2</sub> and (b) Co(CO)<sub>4</sub> on Cu(100) ( $C_{4v}$ ) at  $\beta = 100 \text{ eV}^{-1}$  (T = 116 K). Here, the fully localized limit was used with U = 4.0 eV and J = 0.9 eV. In case of the notations  $0.6z^2 - 0.8(x^2 - y^2)$  and  $0.8z^2 + 0.6(x^2 - y^2)$ , the Co  $d_{z^2}$  and  $d_{x^2-y^2}$  orbitals are mixed after diagonalization of the Co 3*d* subspace.

be further investigated in terms of its Fermi-liquid properties later on). This is an interesting contrast to the bare Co atom on Cu (001), where (at least at relatively large temperature) it is likely the  $d_{z^2}$  orbital which is causing the Kondo properties.<sup>62</sup>

In the case of  $Co(CO)_4/Cu(001)$  in  $C_{4\nu}$  symmetry, we would like to re-emphasize that DFT (using GGA-type functionals) predicts a magnetic moment of  $0.0\mu_B$  on the Co atom. Nonetheless, the spectral function of  $Co(CO)_4/Cu(001)$  obtained from DFT++ [Fig. 2(b)] shows a sharp feature at the Fermi energy for the Co  $3d_{z^2}$  orbital. Based on this observation, there is reason to believe that DFT++ predicts a finite local moment on the Co atom of  $Co(CO)_4/Cu(001)$  in this orbital, which is probably screened due to a Kondo effect and thus leading to the sharp feature in the spectral function, as we will confirm later. Due to the  $C_{4\nu}$  symmetry,  $Co(CO)_4/Cu(001)$  would be a promising candidate for a so-called orbital Kondo effect (as is reported for cobalt-benzene sandwich molecules<sup>63</sup> and for Co adatoms on graphene<sup>64</sup>), because it has two degenerate orbitals ( $d_{xz/yz}$ ). However, the lack of a resonance at the Fermi energy for these orbitals suggests that this is not the case.

To conclude, for  $Co(CO)_2/Cu(001)$  and  $Co(CO)_4/Cu(001)$ ( $C_{4\nu}$ ), we could identify a sharp, Kondo-like feature at the Fermi energy, which gives us the opportunity to investigate the increasing Kondo temperature with an increasing number of CO ligands, for answering the question of how the Kondo effect can be chemically controlled.

#### B. Fermi-liquid properties of Co(CO)<sub>n</sub>/Cu(001)

We study the Fermi-liquid properties to probe the existence of a Kondo effect as suggested by the spectral functions for  $Co(CO)_2$ and for  $Co(CO)_4$  in  $C_{4\nu}$  symmetry.

#### 1. Spin-spin correlation function at high temperatures: Is there a magnetic moment to be screened?

First, we consider the spin-spin correlation functions of the carbonyl cobalt complexes in Fig. 3. Here,  $\chi(\tau)$  is shown at T = 1160 K, because we expect that at this temperature, no Kondo screening takes place, and a finite local moment should be observed.



**FIG. 3.** Spin–spin correlation function  $\chi(\tau)$  of the Co 3*d* orbitals as obtained from PBE++ at  $\beta = 10 \text{ eV}^{-1}$  (*T* = 1160 K) of all carbonyl cobalt complexes on Cu(001) under consideration here. The PBE++ calculation was done with U = 4.0 eV, J = 0.9 eV, and using the fully localized limit for estimating the double-counting correction.

For Co(CO)<sub>2</sub>/Cu(001),  $\chi(\tau)$  of the  $3d_{0.8z^2 + 0.6(x^2-y^2)}$  and  $3d_{yz}$  orbitals drops to a nonzero value as  $\tau \rightarrow \beta/2$ , suggesting a persisting spin density in these orbitals. Most of the magnetic moment observed for Co(CO)<sub>2</sub>/Cu(001) is located in the Co  $3d_{0.8z^2 + 0.6(x^2-y^2)}$  orbital, as suggested by the larger value at  $\chi(\beta/2)$ . This observation suggests that the feature at  $\omega = 0.0 \text{ eV}$  in the spectral function of the Co  $3d_{0.8z^2 + 0.6(x^2-y^2)}$  orbital is indeed a signature of a Kondo effect.

For Co(CO)<sub>4</sub>/Cu(001) ( $C_{4\nu}$ ), only the Co  $3d_{x^2-y^2}$  orbital shows a finite value of  $\chi(\beta/2)$  at T = 1160 K (Fig. 3), which is the same orbital contributing spectral weight at  $\omega = 0.0$  eV in form of a sharp feature. Thus, this orbital is a promising candidate for causing the observed Kondo effect in Co(CO)<sub>4</sub>/Cu(001). For the  $C_{2\nu}$  symmetry, we observe that  $\chi(\beta/2)$  drops to zero for all Co 3*d* orbitals, as for Co(CO)<sub>3</sub>/Cu(001) (see the supplementary material).

To summarize, we were able to identify promising candidates for Kondo-relevant orbitals for  $Co(CO)_2/Cu(001)$  ( $3d_{0.8z^2 + 0.6(x^2-y^2)}$ ) and  $Co(CO)_4/Cu(001)$  in  $C_{4\nu}$  symmetry ( $3d_{x^2-y^2}$ ) from the spectral and spin–spin correlation functions.

#### 2. Temperature dependence of $\chi$ ( $\beta/2$ )

As mentioned at the beginning of Sec. IV, the Fermi-liquid behavior is manifested as a  $T^2$  dependence of  $\chi(\beta/2)$ . In Fig. 4(a), we study this behavior for all Co 3*d* orbitals of Co(CO)<sub>2</sub>/Cu(001) as obtained within the fully localized limit. In all cases,  $\chi(\beta/2)$  drops as *T* is lowered, presumably approaching zero as  $T \rightarrow 0$  K. For the Kondo-relevant orbital  $3d_{0.8z^2} + 0.6(x^2-y^2)$ , one does not observe  $T^2$  dependence (and thus no Fermi-liquid behavior), due to the nonzero intercept with the ordinate. A better agreement within a Fermi-liquid behavior can be observed by shifting the DC value by -1.0 eV from the original FLL value [Fig. 4(b)], which increases the occupation of the  $3d_{0.8z^2} + 0.6(x^2-y^2)$  orbital from 1.31 (FLL) to 1.49 electrons (see Fig. 5).

Here, one observes that the  $3d_{0.8z^2 + 0.6(x^2-y^2)}$  orbital of  $Co(CO)_2/Cu(001)$  behaves quadratically in *T* below T = 150 K. Shifting the DC value by +1.0 eV from the original FLL value, the filling in the  $3d_{0.8z^2 + 0.6(x^2-y^2)}$  orbital reduces to 1.15 electrons, and the values of  $\chi(\beta/2)$  are significantly increased as compared to the results obtained within the FLL. The sensitivity of  $\chi(\beta/2)$  to the shift of the DC value comes from the fact that the occupation on the  $3d_{0.8z^2 + 0.6(x^2-y^2)}$  orbital of  $Co(CO)_2/Cu(001)$  changes from 1.15 (FLL+1 eV) to 1.49 (FLL-1 eV), as shown in Fig. 5. This indicates that the charge fluctuations in this orbital are strongly increased when the DC value is shifted by -1 eV. This leads to the observed transition to the Fermi-liquid regime at roughly T = 150 K, whereas for the other fillings under consideration here, it is not possible to see this transition for the temperature range considered.

In Fig. 4(c), we depict  $\chi(\beta/2)$  as a function of *T* for Co(CO)<sub>4</sub>/Cu(001) ( $C_{4\nu}$ ) as obtained within the fully localized limit.  $\chi(\beta/2)$  of the Kondo-relevant orbital  $(3d_{x^2-y^2})$  behaves as  $T^2$  at temperatures below T = 165 K, pointing to a transition to the Fermi-liquid regime at this temperature. Shifting the DC by  $\pm 1 \text{ eV}$  with respect to the FLL value alters the temperature at which the transition to the Fermi-liquid regime is observed only little [see Fig. 4(d)]. As one can see from Fig. 5, this might be due to the fact that the filling on the  $3d_{x^2-y^2}$  orbital [1.12 (FLL+1 eV) to 1.25



**FIG. 4.** (a)  $\chi(\beta/2)$  as a function of T for all Co 3d orbitals of Co(CO)<sub>2</sub>/ Cu(001) using the fully localized limit for the double-counting correction. (b)  $\chi(\beta/2)$  as a function of *T* for the Co  $3d_{0,8z^2+0.6(x^2-y^2)}$  orbital of Co(CO)<sub>2</sub>/  $3d_{0.8z^2 + 0.6(x^2-y^2)}$  orbital of Co(CO)<sub>2</sub>/Cu(001). Here, the value of the doublecounting was shifted by  $\pm 1.0 \,\text{eV}$  with respect to the FLL value. (c)  $\chi(\beta/2)$  as a function of T for all Co 3d orbitals of  $Co(CO)_4/Cu(001)$  (C<sub>4v</sub>) using the fully localized limit for the double-counting correction. (b)  $\chi(\beta/2)$  as a function of T for the Co  $3d_{x^2-y^2}$  orbital of Co(CO)<sub>4</sub>/Cu(001) (C<sub>4v</sub>). Again, the value of the double-counting was shifted by  $\pm 1.0 \,\text{eV}$  with respect to the FLL value. The fits are quadratic fits of the first two data points. Note that the fits are meant as a guide for the eye to check whether the Fermi-liquid behavior is fulfilled or not.

(FLL-1 eV) electrons] is not affected as strongly as in the case of the  $3d_{0.8z^2 + 0.6(x^2-v^2)}$  orbital in Co(CO)<sub>2</sub>/Cu(001).

#### 3. First Matsubara-frequency rule

We find that a similar behavior of  $Co(CO)_2/Cu(001)$  concerning the Fermi-liquid properties can be found by considering the so-called first Matsubara-frequency rule depicted in Figs. 6(a) and 6(b). The only agreement with the Fermi-liquid behavior can be observed if the DC value is shifted by -1.0 eV,



**FIG. 5.** Co 3*d* fillings of Co(CO)<sub>2</sub> and Co(CO)<sub>4</sub> on Cu(001) as obtained from PBE++ at  $\beta = 100 \text{ eV}^{-1}$  for different values of the double-counting correction.  $E_{\text{DC}}^{\text{FLL}}$  is obtained from Eq. (6).

as indicated by the linear behavior of  $Im\Sigma(\omega_0)$  as  $T \to 0 \text{ K}$  for temperatures below T = 150 K. Thus, both  $\chi(\beta/2)$  and  $Im\Sigma(\omega_0)$ as a function of T lead us to conclude that the  $3d_{0.8z^2} + 0.6(x^2-y^2)$ orbital behaves as a Fermi liquid, but the transition temperature strongly depends on the Co 3*d* filling. Nevertheless, this gives further support to our initial conclusion that the sharp feature observed in the spectral function [Fig. 2(a)] is a signature of a Kondo effect.

For Co(CO)<sub>4</sub>/Cu(001), the first Matsubara-frequency rule [Figs. 6(c) and 6(d)] is also fulfilled for the  $3d_{x^2-y^2}$  orbital at all DC values under consideration here and furthermore affirms the transition to the Fermi-liquid regime roughly below T = 165 K. This gives reason to believe that the observed feature at  $\omega = 0.0$  eV in the spectral function is a true Kondo signature.

#### 4. Discussion

To summarize, we could show that the suggested Kondo-relevant orbitals of  $Co(CO)_2/Cu(001)$  and  $Co(CO)_4/Cu(001)$  ( $C_{4\nu}$ ) display the Fermi-liquid properties expected for a Kondo system. However, for  $Co(CO)_2/Cu(001)$ , the transition temperature, or Kondo temperature, depends delicately on the choice of the double-counting value. Within this work, it is only possible to see a transition to the Fermi-liquid regime (at T = 150 K) if the filling on the Co  $3d_{0.8z^2 + 0.6(x^2-y^2)}$  orbital is increased by shifting the DC value by -1 eV from the FLL value. In the case of  $Co(CO)_4/Cu(001)$  ( $C_{4\nu}$ ), the transition to the Fermi-liquid regime is more robust against changes of the double-counting value, as confirmed unanimously by  $\chi(\beta/2)$  and Im $\Sigma(\omega_0)$  as a function of T. We find that for all DC



**FIG. 6.** (a)  $Im\Sigma(\omega_0)$  as a function of *T* for all Co 3*d* orbitals of  $Co(CO)_2/$ Cu(001) using the fully localized limit. (b)  $Im\Sigma(\omega_0)$  as a function of *T* for the Co  $3d_{0.8t^2 + 0.6(x^2-y^2)}$  orbital of  $Co(CO)_2/$ Cu(001). Here, the value of the doublecounting was shifted by  $\pm 1.0$  eV with respect to the FLL value. (c)  $Im\Sigma(\omega_0)$ as a function of *T* for all Co 3*d* orbitals of  $Co(CO)_4/Cu(001)$  ( $C_{4v}$ ) using the fully localized limit. (d)  $Im\Sigma(\omega_0)$  as a function of *T* for the Co  $3d_{x^2-y^2}$  orbital of  $Co(CO)_4/Cu(001)$  ( $C_{4v}$ ). The fits in (a)–(d) are linear fits as a guide for the eye to check whether the Fermi-liquid behavior is fulfilled or not.

values, the transition to the Fermi-liquid regime of  $Co(CO)_4/Cu(001)$  is at roughly T = 165 K.

Concerning the question of how the Kondo effect is controlled by the number of CO ligands, one is now faced with the problem that the exact double-counting correction for both systems is unknown and that the Kondo temperature of  $Co(CO)_2/Cu(001)$  is only detectable (within the electronic temperatures reached here) if the DC correction is shifted towards smaller values. However, comparing the Fermi-liquid behavior of both systems as obtained from the fully localized limit,  $Co(CO)_4/Cu(001)$  has indeed a larger Kondo temperature than  $Co(CO)_2/Cu(001)$  in qualitative agreement with the experimental data, since in the latter case, the transition to the Fermi-liquid regime occurs at temperatures lower than the ones considered here. Later, we will give an estimation of  $T_K$ based on the hybridization function of the Co  $3d_{0.8z^2} + 0.6(x^2-y^2)$ orbital in  $Co(CO)_2/Cu(001)$ , which confirms this assumption.

### C. Structure dependence of the local moment in $Co(CO)_2/Cu(001)$

In the context of Sec. III, one could ask how strongly the results for  $Co(CO)_2/Cu(100)$  would change for a different molecular structure (see Sec. III for more details about how strongly structural parameters can vary depending on the computational parameters), as this might allow for a deeper insight into structure–property relations for the Kondo effect. To investigate the dependence of the magnetization on the adsorption distance, we show the spin–spin correlation function at T = 1160 K of the Co  $3d_{0.8z^2} + 0.6(x^2-y^2)$ orbital for selected structures (Fig. 7). For the BLYP-D3 and

PBE<sub>fix</sub>-D3-optimized structures,  $\chi(\tau)$  drops faster than for the BLYP-D3+*U*-optimized structure. This points to a stronger screening of the local moment if the adsorption distance is decreased. At the same time as the adsorption distance is increased, the bonding angle  $\phi_{C-Co-C}$  increases, too, and one could expect that the screening also depends on this angle. However, for the structures reported here, we can exclude that the spin-spin correlation function of the Co  $3d_{0.8z^2} + 0.6(x^2-y^2)$  orbital is significantly affected by  $\phi_{C-Co-C}$  (see the supplementary material for further details).



**FIG. 7.** (a) Spin–spin correlation  $\chi(\tau)$  of the Co 3*d* orbitals as obtained from PBE++ at  $\beta = 10 \text{ eV}^{-1}$  (T = 1160 K) of the Kondo-relevant orbital  $3d_{0.8z^2} + 0.6(x^2-y^2)$  in Co(CO)<sub>2</sub>/Cu(001) for different optimized structures. The PBE++ calculations were done with U = 4.0 eV and J = 0.9 eV. (b) Co-surface distance  $d_{\text{Co-surf.}}$ , C–Co–C bonding angle  $\phi_{\text{C-Co-C}}$  and  $3d_{0.8z^2} + 0.6(x^2-y^2)$  filling *n* (PBE++) for Co(CO)<sub>2</sub>/Cu(001) as obtained from different optimized structures.



**FIG. 8.** Projected density of states of the Co 3*d* orbitals  $\left[-\frac{1}{\pi} \text{Im } g_i(\omega)\right]$  and local density of states of the 2*p* orbitals (sum over all 2*p* orbitals) of one of the C atoms. Results obtained from PBE (based on the BLYP-D3+*U* optimized structures).

The occupation of the Co  $3d_{0.8z^2 + 0.6(x^2-y^2)}$  orbital shows a delicate dependence on the structure, similar to its dependence on the DC value as shown for the BLYP-D3+U optimized structure. For this reason, it can be assumed that the transition to the Fermi-liquid regime is similarly affected by changes in the adsorption distance and the C-Co-C bonding angle and, therefore, is very sensitive to the computational parameters with which the molecular adsorbates have been optimized. The structures under study here are likely a particularly challenging case because of their large structural flexibility (as opposed to more rigid phthalocyanines) and because of the challenges associated with describing CO binding to metal surfaces ("CO-puzzle"), as discussed in Sec. III. These results also suggest that the Kondo effect in  $Co(CO)_2/Cu(001)$  might be controllable via external stimuli affecting the adsorption distance (in particular, <sup>-67</sup>). interactions between STM tips and adsorbates<sup>65</sup>

#### V. DFT-BASED ANALYSIS OF THE KONDO EFFECT: WHAT CAN WE LEARN WITHOUT SOLVING THE ANDERSON IMPURITY MODEL?

To gain further insight into the Kondo properties as obtained from solving the Anderson impurity model, it can be helpful to consider the spin-unpolarized PBE electronic structures, as they were used to parametrize the AIM (based on the BLYP-D3+U optimized structures).

In Fig. 8, we show the projected density of states  $\left[-\frac{1}{\pi}\operatorname{Im} g_i(\omega)\right]$  of the Co 3*d* orbitals, as well as the C 2*p* local density of states (LDOS), which is the sum of the projected density of states of the C 2*p* orbitals. They should exhibit peaks at the  $\sigma_2/\sigma_2^*$  and  $\pi/\pi^*$  orbitals of the CO ligands (a schematic representation of the molecular orbital diagram of an isolated CO molecule is provided in the supplementary material). From this, one can learn which of the Co 3*d* orbitals are interacting with the CO ligands, indicated by features in the Co 3*d* PDOS at the same position as the C 2*p* LDOS. This information might be useful to learn more about chemical control of the Kondo effect by increasing the number of CO ligands, as this will affect the coupling of the Co 3*d* orbitals with the rest of the system.

For Co(CO)<sub>2</sub>/Cu(001), the  $3d_{0.8z^2+0.6(x^2-y^2)}$  orbital shows only small features at the C 2*p* LDOS, as a consequence of this orbital interacting only little with the CO ligands (as, e.g., compared to the  $3d_{xz}$  orbital). In contrast, the  $3d_{x^2-y^2}$  orbital in Co(CO)<sub>4</sub> ( $C_{4\nu}$ ) interacts with the CO ligands, as indicated by the features in the PDOS at the position of the  $\sigma_2/\sigma_2^*$  orbitals of CO. This coupling might increase the Kondo temperature, as discussed below in more detail.

Considering the value of the energy-dependent hybridization function (Fig. 9) at the Fermi energy ( $\omega = 0.0 \text{ eV}$ ), one gets a more complete picture of how strongly the Co 3*d* orbitals are coupled with the rest of the system. This value is known from the simplest Kondo model (oneband with a constant hybridization)<sup>50</sup> to be directly connected to the Kondo temperature.

Focusing on the Kondo-relevant orbitals of  $Co(CO)_2/Cu(001)$  $(3d_{0.8z^2+0.6(x^2-y^2)})$  and  $Co(CO)_4/Cu(001)$   $(3d_{x^2-y^2})$ , the energydependent hybridization function is in both cases rather featureless in the range of  $\omega = -1.0 \text{ eV}$  to  $\omega = +1.0 \text{ eV}$ . As shown in Table II, the value at  $\omega = 0.0 \text{ eV}$  for the  $3d_{x^2-y^2}$  orbital of  $Co(CO)_4/Cu(001)$ is about four times larger than for the  $3d_{0.8z^2+0.6(x^2-y^2)}$  orbital in  $Co(CO)_2/Cu(001)$ . This supports our assumption that the interaction of the  $3d_{x^2-y^2}$  orbital of  $Co(CO)_4$   $(C_{4\nu})$  with the  $CO \sigma_2/\sigma_2^*$ orbitals increases the coupling at the Fermi energy, which in turn results in a larger Kondo temperature  $(T_K \approx 165 \text{ K})$  as discussed in Sec. IV. Assuming that the Kondo temperature for  $Co(CO)_2/Cu(001)$ is lowered by the same factor as the hybridization at the Fermi energy of the  $3d_{0.8z^2+0.6(x^2-y^2)}$  orbital, one would expect it to be



**FIG. 9.** Imaginary part of the Co 3*d* hybridization functions of (a)  $Co(CO)_2$  and (b)  $Co(CO)_4$  ( $C_{4v}$ ) on Cu(100) as obtained form PBE (based on the BLYP-D3+*U* optimized structures).

J. Appl. Phys. **125**, 142910 (2019); doi: 10.1063/1.5079518 Published under license by AIP Publishing.

<b>TABLE II.</b> Hybridization in eV at the Fermi energy $-Im\Delta(0 \text{ eV})$ as obtained from
spin-unpolarized PBE (based on the BLYP-D3+ $U$ optimized structures) for different
carbonyl cobalt complexes on Cu(001). For Co(CO) <sub>2</sub> , the orbitals are labeled
according to their largest contribution after diagonalizing the Co 3d subspace.

	$d_{x^2-y^2}$	d <sub>xz</sub>	d <sub>xy</sub>	d <sub>yz</sub>	$d_{z^2}$
$Co(CO)_2$	0.24	1.48	0.24	0.1	0.33
$Co(CO)_4$ ( $C_{4\nu}$ )	1.31	1.91	5.87	1.91	0.33

around  $T_{\rm K} \approx 41$  K. The lowest electronic temperature under consideration in the DFT++ calculations as discussed in Sec. IV was T = 46.4 K, which could explain why within the fully localized limit, we were not able to reach the transition to the Fermi-liquid regime. According to Sec. IV C, the decreased adsorption distance of Co(CO)<sub>2</sub>/Cu(001) in, e.g., the BLYP-D3-optimized structure, shows a hybridization of the Kondo-relevant orbital at  $\omega = 0.0$  eV of 0.52 eV. Compared with Im $\Delta$ (0 eV) = 0.33 eV obtained for the BLYP-D3+U structure, the Kondo temperature would, therefore, probably shift towards larger values for this structure.

#### VI. CONCLUSION

The chemical and mechanical control of the Kondo effect in molecular adsorbates is an intriguing subject, which promises insight into strong electron correlation. We have studied such control exploring experimentally characterized cobalt carbonyl complexes on Cu(001) from a theoretical point of view, employing both DFT++ approaches for a full description of correlation and DFT-derived properties for a conceptual understanding of structure-property trends. We find that it is indeed possible to optimize structures with DFT (employing BLYP-D3+U) whose Fermi-liquid properties are compatible with the experimentally observed trend of larger Kondo temperatures with increasing number of ligands for the di- and tetracarbonyl complex (constraining the latter to  $C_{4\nu}$  symmetry). We can trace back this behavior to an increased hybridization at the Fermi energy, which correlates with a stronger interaction of the Kondo-relevant 3d orbital with the CO ligands for the tetracarbonyl. This Kondo-relevant orbital is  $d_{x^2-y^2}$  in both cases, with a strong admixture of  $d_{z^2}$  for the dicarbonyl system. It would be interesting to compare these data with newly developed approaches, in which a general projection scheme allows for extending the correlated impurity from the cobalt 3d orbitals to molecular orbitals which include part of the CO ligands.<sup>6</sup>

Our data also point to the challenges such systems pose for present-day first-principles electronic-structure methods: The structural flexibility of cobalt carbonyl complexes, along with the known difficulty of describing direct carbonyl-metal binding by presentday DFT, implies that predictive modeling of their Kondo properties is virtually impossible. In particular, no atomistic structure could be obtained for the tricarbonyl, which is compatible with the experimentally observed Kondo effect (and with the lack of threefold symmetry suggested by STM data). Furthermore, all DFT protocols employed here suggest that for the tetracarbonyl, a  $C_{2\nu}$ -symmetric structure is by at least 30 kJ/mol more stable than a  $C_{4\nu}$ -symmetric one, yet only for the latter can we obtain Fermi-liquid properties consistent with the experimentally observed Kondo effect. This suggests that the fourfold-symmetric structure observed in the STM results from the intrinsic symmetry of the molecule rather than from a rotational process. This discrepancy might result from the deficiencies of present-day DFT or from a kinetic stabilization of the  $C_{4\nu}$ -symmetric structure in the experiment. This is an example of employing spectroscopic data rather than solely total energies for identifying molecular structures compatible with the experiment, as also done, for example, in theoretical EXAFS studies.<sup>70</sup> For systems with a less pronounced structural flexibility, such as metal phthalocyanines, it is likely that available first-principles methods are more reliable at present. On the upside, the strong dependence of Kondo properties on structural parameters suggested by our data could imply that these Kondo properties can be controlled mechanically, particularly by interactions with an STM tip.

#### SUPPLEMENTARY MATERIAL

See the supplementary material for further details on atomistic structures, on spectral properties, and spin–spin correlation data of  $Co(CO)_3$  on Cu(001) and  $Co(CO)_2$  on Cu(001) ( $C_{2\nu}$ ), on molecular orbitals of the CO molecule, and on the angular dependence of spin–spin correlation and hybridization function for  $Co(CO)_2$  on Cu(001).

#### ACKNOWLEDGMENTS

The authors acknowledge the high-performance-computing team of the Regional High-Performance Computing Center at the University of Hamburg and the North-German Supercomputing Alliance (HLRN) for technical support and computational resources, and the DFG for financial support via SFB 668. Furthermore, the authors are grateful to Michael Karolak for valuable discussions.



**FIG. 10.** Density of states of the 4s, 4*p*, and 3*d* orbitals of one of the Cu surface atoms. Results obtained from PBE.



**FIG. 11.** Co 3*d* hybridization functions (imaginary part) as obtained from an isolated (optimized) Co on Cu(001), from an isolated Co(CO)<sub>4</sub> molecule as obtained by removing the surface atoms from the optimized Co(CO)<sub>4</sub> / Cu(001), and from Co(CO)<sub>4</sub> on Cu(001). Hybridization functions as obtained from PBE, based on BLYP-D3+*U* optimized structures. The hybridization functions of an isolated Co on Cu(001) agrees qualitatively with the ones reported in Ref. 72.

#### APPENDIX: THE EFFECT OF THE CO-SURFACE INTERACTION ON THE Co 3*d* HYBRIDIZATION FUNCTION

It would be interesting to study the contribution to the hybridization function of the Co 3*d* orbitals coming from the CO ligands directly, or indirectly as caused by the ligands being coupled to the Cu(001) surface. We try to make a step towards answering this question using the example of Co(CO)<sub>4</sub>/Cu(001) in  $C_{4\nu}$  symmetry. For this purpose, we compare the hybridization functions in Fig. 11 of all Co 3*d* orbitals as obtained from an isolated Co on Cu(001), for an isolated Co(CO)<sub>4</sub> molecule (no surface), and for Co(CO)<sub>4</sub> on Cu(001). For the isolated molecule, we start from the optimized system and remove all Cu atoms, in order to see the contribution to the hybridization function of the CO ligands only. While this analysis neglects the effects of the surface on the CO ligands, for example, which may in turn affect the way these ligands contribute to the hybridization of the Co orbitals, we do expect an elucidating qualitative picture of the relative importance of ligands and surfaces.

Note that the plots in Fig. 11 are differently scaled on the *y*-axis. The hybridization functions of all Co 3*d* orbitals for an isolated Co atom on Cu(001) are rather small and featureless. In all cases, however, there is a small bump at roughly E = -1.8 eV to E = -2.5 eV, which comes from the increased DOS of the Cu surface at this energy (as shown in Fig. 10).

Considering the Co 3d hybridization functions (Fig. 11) of the isolated molecule should give an impression of the contribution of the ligands to the hybridization of the Co 3d orbitals in the full

system. For the isolated molecule, the hybridization functions of all 3*d* orbitals (except for the  $d_{x^2-y^2}$  orbital) exhibit a sharp feature close to the Fermi energy, which is the reason for the hybridization function of these orbitals being increased in the vicinity of  $\omega = 0.0 \text{ eV}$  for the full system  $[\text{Co}(\text{CO})_4/\text{Cu}(001)]^{71}$  compared with an isolated Co atom on Cu(001). We believe that this enhancement is directly induced by the hybridization with the ligands. For the  $d_{x^2-y^2}$  orbital, the hybridization at E = -1.8 eV is significantly increased compared to Co/Cu(001), although for the isolated molecule, we do not observe a peak in  $\Delta(\omega)$  at this energy. Thus, we conclude that the increased hybridization is indirectly caused by the CO ligands, as they couple strongly not only to the Co  $d_{x^2-y^2}$  orbital (as pointed out in Sec. V) but also to the Cu(001) surface.

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