Structural and magnetic properties of FeMn\textsubscript{x} chains (x = 1–6) supported on Cu\textsubscript{2}N/Cu (100)

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Heterogeneous atomic magnetic chains are built by atom manipulation on a Cu\textsubscript{2}N/Cu (100) substrate. Their magnetic properties are studied and rationalized by a combined scanning tunneling microscopy (STM) and density functional theory (DFT) work completed by model Hamiltonian studies. The chains are built using Fe and Mn atoms on top of the Cu atoms along the N rows of the Cu\textsubscript{2}N surface. Here, we present results for FeMn\textsubscript{x} chains (x = 1–6) emphasizing the evolution of the geometrical, electronic, and magnetic properties with chain size. By fitting our results to a Heisenberg Hamiltonian we have studied the exchange-coupling matrix elements J for different chains. For the shorter chains, x ≤ 2, we have included spin-orbit effects in the DFT calculations, extracting the magnetic anisotropy energy. Our results are also fitted to a simple anisotropic spin Hamiltonian and we have extracted values for the longitudinal-anisotropy D and transversal-anisotropy E constants. These parameters together with the values for J allow us to compute the magnetic excitation energies of the system and to compare them with the experimental data.

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I. INTRODUCTION

Bottom-up approaches give access to systems of very reduced dimensionality with unique physical properties. Among these systems, chains of a single-atom cross section are of great interest [1]. When the chains are formed by magnetic atoms, spin-spin correlations can come into place leading to new phenomena and applications [2]. Building structures from their atomic constituents can be achieved by the atomic-manipulation capabilities of the scanning tunneling microscope (STM). Magnetic atoms have been positioned one by one at different distances and with different arrangements on a variety of substrates [3–7]. Spin-polarized STM [8] or inelastic electron tunneling spectra (IETS) [9,10] have granted us a detailed vision of the magnetic mechanisms at play on the atomic scale.

Recent developments in density-functional theory (DFT) have also permitted us to attain a deep understanding of the phenomena revealed by the above experiments. Due to their experimental interest, Mn chains are amongst the most studied ones [11–16]. Fe chains are also well known both experimentally and theoretically [4,5,15,17–21]. The work by Lin and Jones [13,22] on Fe, Co, and Mn atoms on Cu\textsubscript{2}N/Cu (100) reveals that atomic spins maintain their nominal values on the surface (S\textsubscript{Fe} = 2, S\textsubscript{Mn} = 5/2, and S\textsubscript{Cu} = 3/2), showing the interest of using Cu\textsubscript{2}N to preserve much of the magnetic identity of transition-metal (TM) atoms. Studies of Fe chains [17] and of Mn [12] reveals that close-packed TM chains on Cu\textsubscript{2}N/Cu (100) couple antiferromagnetically due to a N-mediated superexchange mechanism, largely explaining experimental findings [5,22].

In the present work, we report on a different type of magnetic atomic chain. These chains are heterogeneous, including two types of magnetic atoms, Fe and Mn, on a Cu\textsubscript{2}N/Cu (100) substrate. An initial account of the results has been given in a separate publication [23]. The experimental study is based on STM and IETS results of FeMn\textsubscript{x} (x = 1–6) and confirms that, as in the previous cases, the magnetic ordering along the chain is antiferromagnetic. The different anisotropy of Fe and Mn on Cu\textsubscript{2}N/Cu (100) leads to two possible orientations of the magnetic moments (along the chain and out of plane). Contrary to the case of homogeneous chains, this may indicate a noncollinear arrangement. When x is an odd number, a simple-minded evaluation of the total spin yields 1/2, which is compatible with the appearance of a Kondo feature at zero bias [23]. However, when x is even, the expected spin is 2 and, correspondingly, no Kondo peak is observed. In the present work, we perform extensive calculations and compare them with the experiments. The comparison permits us to conclude on the spin arrangement (it is collinear along the chain) and on the behavior of the exchange coupling between the two different atomic species of the chain in good agreement with the experimental observations. The antiferromagnetic coupling is confirmed and traced back to the N-mediated superexchange mechanism as in the homogeneous chains. Finally, the obtention of model Hamiltonians to study the magnetic structure of these chains is discussed within the framework of DFT calculations [18,24].

II. EXPERIMENTAL METHOD

Experiments were performed in an ultrahigh vacuum low-temperature STM at a base temperature of 0.5 K as has been
partially reported in Ref. [23]. Differential conductance was used as a local spectroscopic tool that gives information via inelastic electron tunneling (IETS) [9,10]. The differential conductance was directly measured using lock-in detection with 72-µV rms modulation at 691 Hz to the sample bias voltage $V$.

The Cu (100) surface was cleaned by Ar sputtering and then annealed up to 850 K. After having big terraces of Cu(100) crystal, a monolayer of Cu$_2$N was formed as a decoupling layer by nitrogen sputtering and post-annealing at 600 K. Single Fe and Mn atoms were deposited onto the pre-cooled surface. Pt-Ir tips were prepared by sputter-anneal cycles and coated with copper in vacuo by soft indentations into the Cu bridges. The tip status was monitored through STM images and controlled to manipulate the atoms. All atomic chains were built using vertical atomic manipulation. After identifying a given adatom by its spectroscopic features, we picked it up by voltage pulsing, dropped it off on a nitrogen site and hop it to a copper site. We build the chain atom by atom in a close-packed configuration to ensure an AF coupled spin chain. By doing so, all different kinds of FeMn$_x$ ($x = 1$–10) chains were constructed.

III. THEORETICAL METHOD

Ab initio calculations are performed within the density-functional theory (DFT) framework as implemented in the VASP code [26]. We expand the wave functions using a plane-wave basis set with a cutoff energy of 300 eV. Core electrons are treated within the projector augmented wave method [27,28]. The PBE form of the generalized gradient approximation is used for the exchange and correlation functional [29].

The system consists of a Cu(100) surface covered by a Cu$_2$N layer. To model the surface, we use a slab geometry. For Mn, they find $U = 4.9$ eV when Mn is sitting ontop a surface Cu atom, while it is reduced to $U = 3.9$ eV ontop on CuN substrate. Rudenko et al. [12] take an effective value $U - J = 5$ eV, where $J$ is the intra-atomic exchange coupling. Nicklas et al. [17] do not take any value of $U$ for Fe. As Lin and Jones [13] show, the actual value of $U$ greatly affects the computed exchange couplings, indeed, they find a difference of a factor of 2 between their calculations with $U = 7$ eV and without $U$, finding DFT+$U$ results in better agreement with the experiment.

In this work, we have used the GGA+$U$ method of Dudarev [31] with a $U_{eff} = U - J$ of 1 eV for Fe and 4 eV for Mn. The chosen values correspond to roughly subtracting $J$ to 1 eV for Fe and $U = 7$ eV as computed by Lin and Jones [13] for Mn (Fe) ontop a Cu atom. As we will see, the values of $U_{eff}$ become critical for the determination of the exchange couplings. The effect of different values for $U_{eff}$ is discussed below.

To rationalize our results, we have fitted them to a Heisenberg Hamiltonian in the form

$$H = -\sum_{i,j>i} J_{ij} S_i \cdot S_j, \quad (1)$$

where $J_{ij}$ are the exchange couplings between spins $S_i$ and $S_j$. Evaluating different magnetic configurations we have been able to extract the $J_{ij}$ values by fitting the DFT energies to the Heisenberg Hamiltonian, Eq. (1).

For the shorter chains (FeMn and FeMn$_2$), we have included spin-orbit coupling (SOC) [32,33] into our scalar-relativistic Hamiltonian. We have considered a simple anisotropic spin Hamiltonian, in the form

$$H = DS^2 + E(S^2 - S^2_{\parallel}). \quad (2)$$

Here, $i$, $j$, and $k$ are such that $i$ is the computed magnetic axis, while $j$ and $k$ are two orthogonal directions in the plane perpendicular to $i$ (hard-plane) in principle, different from the surface directions. The energy of the system has been self-consistently calculated including SOC for the different orientations $i$, $j$, and $k$ of the magnetization axis and, from these energies, the values for $D$ and $E$ in Eq. (2) have been fitted using the evaluated magnetic moments.

IV. RESULTS

A. Geometries and energetics

Constant current images were obtained between atom manipulation sequences. This permitted us to have precise knowledge of the atomic arrangement. Unfortunately, the structure of the tip that is optimal for atom manipulation is not necessarily good for image production and the obtained images do not contain much information, as can be seen in Fig. 1. In the inset, we show the configuration as inferred from the atomic manipulation procedure that corresponds to a FeMn$_3$ chain.

The calculation yields precise insight on the actual geometries. In Fig. 2, we show the FeMn$_2$ geometry as an example. As we can see in the side view, after deposition of the TM atoms the final chain includes the N atoms in between them. These atoms are lifted from the surface. N atoms at the edge of the chain, like N$_1$, are moved upwards by $\approx 0.7$ Å, while N atoms in between TM atoms, like N$_2$, are lifted by $\approx 1.6$ Å with respect to their unrelaxed positions in the bare CuN$_2$ surface. Therefore we can assume that the final chains have the form FeMn$_x$N$_{x+2}$. Of course, this is just a conventional choice to
identify the final chain, since other atoms in the surface are also significantly disturbed and could be considered as parts of the chain, like CuS1 and CuS2.

It is interesting to study the evolution of the geometry of the chains as their size increases. In Table I, we show the distance between TM atoms within the chain for all considered sizes. We observe that these distances are bigger than the unrelaxed value (3.65 Å corresponding to the PBE lattice parameter of Cu due to the arrangement of the chain on the surface). For example, $d_{[Fe-Mn_1]}$ for FeMn is 3.81 Å, 4% bigger than the unrelaxed value. This distance tends to decrease as the chain size is increased, reaching 3.73 Å for FeMn6. The same behavior can be observed for other positions in the chain. Another interesting distance to track involves the Cu atoms just underneath Fe and Mn atoms. As can be seen in the side view of Fig. 2, these Cu atoms (CuFe, Cu1, Cu2) are pushed downwards by around 0.4 Å. For FeMn, the Fe-CuFe distance is very close to the value for the isolated atom (2.37 Å versus 2.38 Å). However, it increases for longer chains, with a value of 2.40 Å for FeMn6. The same behavior is observed for the Mn atoms, again close to the single adsorbate for the case of FeMn (2.50 Å versus 2.47 Å), and it increases for longer chains reaching 2.56 Å for FeMn6. Hence, as the number $x$ of Mn atoms increases, we find that the intra-chain distances diminish, see Table I, while the distance between the chain atom and the top Cu ones increases. Both behaviors can be rationalized as stress built up as the chain increases its size. Indeed, the elongation of these distances can be easily identified with some destabilization of the chains as length is increased.

To check for the stability of the chains we have studied the chain’s energetics. First, we start by analyzing the atomization energy per TM atom, $E_{at}[FeMn_x]$, defined from

$$E_{at}[FeMn_x] = -\frac{1}{x+1} \left[ E[FeMn_x] - E[Fe] - x E[Mn] - E_{Cu2N} \right],$$

where $E[FeMn_x]$ is the total energy of the system; $E[Fe]$ ($E[Mn]$) is the energy of a gas-phase Fe (Mn) atom; and $E_{Cu2N}$ is the energy of the [3 × (x + 3)] unit cell of the bare surface. The atomization energy per TM atom is a measure of how much average energy per atom one needs to give to the adsorbed chain to separate it in its constituent Fe and Mn gas-phase atoms and the pristine Cu2N/Cu(100) substrate. Figure 3 shows the atomization energy per TM atom as a function of the number of Mn atoms, $x$. The atomization energy per TM atom tends to decrease as the chain size increases, implying that long chains become energetically less favorable.

We can extract more information by looking at another quantity that reflects better the experimental procedure to construct the chain. Let us remember that the chains are constructed by STM manipulation. First, Fe and Mn atoms are deposited on the surface, and then they are moved with the STM tip to form the desired structure. To account for this procedure, we can define a formation energy where the reference of energy is the one of the deposited TM atom, and then to study the gain in energy by adding one more atom to the chain. The definition that we used for the formation energy is given by

$$E_F[FeMn_x] = -[E_{at}[FeMn_x] - E_{at}[Fe] - E_{at}[Mn]],$$

where we use the atomization energies defined in Eq. (3). From $E_F$, we can define the gain in energy by adding one more TM
TABLE I. Distances between the different TM atoms for the six considered chains (in angstroms). Note that the unrelaxed value would be 3.65 Å in agreement with the DFT lattice parameter of the Cu substrate.

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<td>3.75</td>
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<td>3.74</td>
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atom, $\Delta_x E_F$ [34], as

$$\Delta_x E_F = E_F[\text{Fe}\text{Mn}_x] - E_F[\text{Fe}\text{Mn}_{x-1}]$$

$$= -[E_{al}[\text{Fe}\text{Mn}_x] - E_{al}[\text{Fe}\text{Mn}_{x-1}] - E_{al}[\text{Mn}]].$$  (5)

This quantity, $\Delta_x E_F$, defines the energy gained by adding a Mn atom to an existing chain starting from an Fe atom, which mimics the experimental procedure to build the system. The calculated values of $\Delta_x E_F$ are always positive indicating that building the chains is an energetically favorable process. The energy gain is modest ($\approx$0.4 eV), and decreases by 37% from FeMn to FeMn6 (Fig. 3). Experimentally, the longest chains contained nine Mn atoms [23]. Our results seem to suggest that longer chains will be difficult to form due to the accumulated stress imposed by the surface lattice constant and the Fe–Mn distance.

B. Electronic and magnetic properties and charge transfer

The magnetic properties are one of the main motivations for the study of these systems. They largely stem from the electronic structure of the atomic chains on the surface. An analysis of the electronic structure by projection onto atomic orbitals have been shown in Ref. [23]. The magnetic properties were shown to be given by the population of Fe and Mn d orbitals. In the case of Fe, the $d_x^2$ orbital was basically filled (where $z$ is the direction perpendicular to the surface in the present case), not contributing to the atomic magnetic moment. The Mn atoms show a larger spin polarization due to the complete polarization of its $d$ shell after inclusion in the FeMn$_x$ chain. The exchange splitting of $d$ bands is the smallest for Fe, but still it is roughly 5 eV, showing how robust magnetism is in these chains. The electronic structure near the Fermi energy is largely due to the Cu and N states and with negligible magnetic polarization.

It is interesting to inspect the total magnetic moment of the chains shown in Table II. We can see how the magnetic couplings between Fe-Mn and Mn-Mn atoms are antiferromagnetic (AF) in the most stable configuration. In addition, the magnetic moment of atomic Fe on the surface is 3.35 $\mu_B$ (formally $S = 2$), while the value for Mn is 4.80 $\mu_B$ (formally $S = \frac{5}{2}$). Therefore formally, when adding Mn atoms to a Fe one, we would get systems with $S = \frac{x}{2}$ for an odd number of Mn atoms, $x$, and $S = 2$ for an even number. In the computed results for the total magnetic moment in the cell ($\mu_B$), we observe the expected even-odd behavior, with values of 1.30–1.43 Bohr magnetons, $\mu_B$, for $x$ odd, and 3.68–3.86 $\mu_B$ for the $x$ even. Values of 1.30–1.43 $\mu_B$ correspond well with the simple result $S = \frac{x}{2}$. The difference from $1 \mu_B$ has two different sources that are present in these calculations. On the one hand, the TM atoms present fractional occupancies that lead to noninteger multiples of $\mu_B$. On the other hand, these calculations are mean-field approximations to the difficult solutions of correlated AF ground states. As a consequence, the mean-field solution averages over the possible atomic magnetic moments found in the multiple spin configurations of the correct AF ground state.

The AF coupling between the TM atoms is mediated by the N atoms via a superexchange mechanism as previously shown for Mn [12] and Fe chains [17]. To illustrate the superexchange interaction, we show the spin-difference-density of the FeMn$_3$ chain in Fig. 4. The spin polarization of the intercalated N atoms adopts the expected form for superexchange, with induced spin polarization within the atom, but net spin close to zero. Indeed, the bigger values for the induced magnetic moments of N atoms are very small, for example 0.11 $\mu_B$ for N$_1$ and 0.08 $\mu_B$ for N$_2$. Superexchange also leads to a change in the atomic angles: the Fe-N-Mn and Mn-N-Mn angles tend to approach 180° maximizing the AF interactions [12,17].

The magnetic moments tend to increase with increasing chain size, which is consistent with the previously mentioned progressive separation of the chain from the surface. The magnetic moments of the Mn atoms within a given chain are...
quite constant, with the exception of the final atom, which is
≈4% bigger. The reason is the lower coordination of the edge
atom, reflected in the slightly longer distance with the Cu atom
below and the longer distance with the adjacent N atoms.

The analysis of charge transfer in the system yields that
each TM atom in the chain loses around one s electron to form
the bond with N atoms in the chain and the Cu atom underneath
the TM atom. The d charge is similar to the atomic case, i.e.,
there are six electrons in the d manifold for Fe, and five for
Mn, in agreement with the computed magnetic moments.

TABLE II. Magnetic moments (in $\mu_B$) of the TM atoms in the chains. Values for isolated Fe and Mn atoms on the surface are included for comparison. The values have been obtained by performing a Bader analysis [30]. The total magnetic moment $\mu_T$ of the calculation cell also includes contributions from nearby copper atoms that spin polarize, hence $\mu_T$ slightly differs from the sum of the magnetic moments of Fe and Mn.

<table>
<thead>
<tr>
<th>Atom</th>
<th>$\mu_T$ (eV)</th>
<th>$\mu_{[Fe]}$ (eV)</th>
<th>$\mu_{[Mn_1]}$ (eV)</th>
<th>$\mu_{[Mn_2]}$ (eV)</th>
<th>$\mu_{[Mn_3]}$ (eV)</th>
<th>$\mu_{[Mn_4]}$ (eV)</th>
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<tr>
<td>Fe</td>
<td>4.97</td>
<td>4.80</td>
<td>4.80</td>
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<td>Mn</td>
<td>3.82</td>
<td>3.35</td>
<td>3.78</td>
<td>3.23</td>
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<tr>
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<td>−4.65</td>
<td>−4.58</td>
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<tr>
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<td>−4.56</td>
<td>4.56</td>
<td>−4.73</td>
<td>4.73</td>
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<td>−4.56</td>
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<tr>
<td>FeMn$_6$</td>
<td>3.86</td>
<td>3.23</td>
<td>−4.57</td>
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</table>

C. Exchange coupling constants

We have fitted our results to the Heisenberg Hamiltonian
shown in Eq. (1). In order to do so, we have calculated different
spin solutions corresponding to different magnetic ordering
among the TM atoms in the chain. In all calculations, we have
used the geometry of the most stable configuration, which is
always the AF case shown in Table II. For the atomic values of
$S$, in Eq. (1), we had the choice of either using the formal value
($S = 2$ for Fe, $S = 5/2$ for Mn), or taking the computed value
from the DFT calculation since for a single atom $S_z$ is a good
quantum number. We have opted for the later option, which
gives values for the exchange coupling constants $J$’s, which
are between 15% and 20% higher. We can justify this choice
because the DFT values reflect the local magnetic moment
that interacts via other atoms (superexchange) with the local
magnetic moment of the next neighbor in the calculation. The
results are shown in Table III. As expected, $J$’s involving first
neighbors are negative, which indicate an AF coupling. Second
neighbor $J$’s are positive, indicating a ferromagnetic coupling,
which further stabilizes the global AF solution. Further order
$J$’s are smaller, and its omission in the fit only implies a small
error.

The convergence with the number of coupled atoms con-
sidered in the Heisenberg chain, Eq. (1), is very fast. We have
done a systematic study for FeMn$_4$ where different number
of neighbors are included in solving Eq. (1). Considering
just first neighbors introduces a maximum error of 0.2 meV.
Considering first and second neighbors the error is less than
0.1 meV. These calculations show that the effective interactions
in these spin chains are very short range and first-neighbors
truncation is indeed an accurate approximation.

Analyzing the evaluated exchange couplings, $J$, we observe
that the biggest value is obtained for the first-neighbors
interaction between Fe and Mn$_1$. Thus the AF coupling
between Fe and Mn is stronger than the one between Mn-Mn.
FeMn is a special case since it involves two edge atoms, the
resulting AF coupling is the strongest one. Curiously, the
exchange coupling between Fe and Mn$_1$ ($J_{12}$) presents a sharp
drop for FeMn$_2$ to start increasing again for FeMn$_3$ and FeMn$_4$.
The coupling between Mn$_1$ and Mn$_2$ ($J_{23}$) presents the same
behavior, it shows a minimum for FeMn$_3$. Similar behavior is
obtained for pure Mn chains by Rudenko et al. [12]. These
minima in the exchange couplings seem to be independent
from the atomic geometry, where the behavior with chain
length is monotonous, and with the atomic magnetic moments.
It is probably due to the sudden appearance of an extra neighbor
that symmetrizes the interactions on the central atom of the
chain.

As Rudenko et al. [12] and Lin and Jones [13] note, there
is a dependence of the values of the interatomic exchange
coupling on the chosen value of $U_{eff} = U - J$ for the GGA+U
approximation. Figure 5 shows the dependence of $J_{12}$ for
FeMn on $U_{eff}$. We find that as the value of $U_{eff}$ decreases,
$|J_{12}|$ increases. For the values of the present study [13] ($U_{eff}$
[Fe] = 1 eV and $U_{eff}$ [Mn] = 4 eV) $J_{12} = -10.25$ meV. If
$U_{eff}$ [Fe] is reduced by 1 eV, this value changes to $-11.60$
meV. When $U_{eff}$ [Mn] is reduced by 1 eV the effect is more
pronounced, obtaining a value of $-16.44$ meV (60% increase
in absolute value). These values give us an interval of exchange
couplings that will be compared with our experimental results
in the following sections.

![FIG. 4. Spin density of the FeMn$_3$ chain. Red (yellow) indicates majority (minority) spin areas. The chosen isovalue is 0.01 eV/Å$^3$. The color code of the atoms is the same of Fig. 2.](image-url)
approximations. Using GGA MAE calculations are very robust against different DFT cell size [37] and present a negligible dependence. Moreover, similar results to the strictly LDA calculations by Shick with a value of 0.15 meV in good agreement with Ref. [35].

For Mn, the MAE is much smaller and larger values of the energy as compared with previous calculations [35,36]. The transversal direction show the easy axis directions, where

\[ E = \frac{E_j - E_k}{S(2S - 1)} \]

where \( E_i \), \( E_j \), and \( E_k \) are the magnetic anisotropy energies when the spin, \( S \), is aligned along the easy or hard axis \( i \) and the two transversal axes \( j \) and \( k \), see Table IV.

For Fe, the minimum of MAE lies along the N direction, \( i = y \), with a value of 1.88 meV in good agreement with previous calculations [35,36]. The transversal direction show larger values of the energy as compared with previous calculations [35,36]. For Mn, the MAE is much smaller and with a value of 0.15 meV in good agreement with Ref. [35].

Our results have been tested against convergence with unit cell size [37] and present a negligible dependence. Moreover, MAE calculations are very robust against different DFT approximations. Using GGA+U, Barral et al. [36] obtained similar results to the strictly LDA calculations by Shick et al. [35]. Panda et al. [38] obtain a small change in D when they use an extraordinarily large value of \( U \) (from \( D = -0.48 \) meV for \( U = 0 \) to \( D = -0.67 \) meV for \( U = 6 \) eV). Hence, as Panda et al. [38] show, improved calculations of MAE need to go beyond DFT.

The calculation of \( D \) and \( E \) differ from reported values. Our calculations compared better with the values of Ref. [35] if the nominal magnetic moments are used. Indeed, we obtain \( D = -0.44 \) meV and \( E = 0.19 \) meV using \( S = 2 \) for Fe and \( D = -0.67 \) meV, \( E = 0.29 \) meV for the computed magnetic moments. These values are obtained if we assume an easy axis along the \( y \) direction as found experimentally [22]. For Mn the agreement with Ref. [35] improves probably because the DFT magnetic moment agrees with the nominal one. However, Barral et al. [36] report on a different spin direction for Mn. As these last authors discuss, the MAE values for Mn on Cu2N are very small and prone to numerical inaccuracy. These results support the idea that MAE calculations require post-DFT treatments such as the very recent ones of Ref. [38].

Taking into account that experimentally Mn and Fe display perpendicular easy axes [22], one may wonder if the SOC might induce a noncollinear alignment of the magnetic moments in the chains. We have tested that possibility for the shorter chains by performing an explicit noncollinear calculation including SOC [33]. Due to the small energies involved in the calculation, the convergence of noncollinear solutions is very challenging. We have been able to stabilize a noncollinear configuration for FeMn where the magnetic moment of Fe is along the chain, while the moment for Mn forms an angle of 27° with the \( y \) axis. However, this solution is 3.79 meV higher in energy than the collinear solution. For longer chains, we have not been able to stabilize any noncollinear solution. Therefore we have just considered collinear configurations for the rest of our calculations.

The larger MAE of Fe forces the full-chain magnetic axis to align with the Fe anisotropy axis. This is indeed

### Table III. Values for the exchange coupling constants (in meV) obtained by fitting Eq. (1) to the GGA+U total energies.

<table>
<thead>
<tr>
<th></th>
<th>First neighbors</th>
<th>Second neighbors</th>
<th>Third neighbors</th>
<th>Fourth</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>( J_{12} )</td>
<td>( J_{13} )</td>
<td>( J_{14} )</td>
<td>( J_{15} )</td>
</tr>
<tr>
<td>FeMn</td>
<td>10.25</td>
<td>−6.51</td>
<td>0.97</td>
<td>0.73</td>
</tr>
<tr>
<td>FeMn(_2)</td>
<td>−6.51</td>
<td>−2.09</td>
<td>1.14</td>
<td>0.05</td>
</tr>
<tr>
<td>FeMn(_3)</td>
<td>−7.30</td>
<td>−1.97</td>
<td>0.10</td>
<td>−0.36</td>
</tr>
<tr>
<td>FeMn(_4)</td>
<td>−7.96</td>
<td>−2.69</td>
<td>−2.10</td>
<td>−0.02</td>
</tr>
</tbody>
</table>

### Table IV. Magnetic anisotropy energies (MAE) (in meV). We also show the easy axis directions, where \( x \) is the direction in plane and perpendicular to the chain, \( y \) along the chain, and \( z \) perpendicular to the surface.

<table>
<thead>
<tr>
<th>Material</th>
<th>( E_x - E_y )</th>
<th>( E_z - E_y )</th>
</tr>
</thead>
<tbody>
<tr>
<td>Fe</td>
<td>0.74</td>
<td>1.88</td>
</tr>
<tr>
<td>Mn</td>
<td>−0.15</td>
<td>−0.02</td>
</tr>
<tr>
<td>FeMn</td>
<td>0.75</td>
<td>1.51</td>
</tr>
<tr>
<td>FeMn(_2)</td>
<td>0.48</td>
<td>1.46</td>
</tr>
</tbody>
</table>

FIG. 5. Evolution of \( J_{12} \) for FeMn with respect to the variation of \( U_{\text{eff}} \). We have changed one of the values of \( U \) while keeping constant the other one.
seen in Table IV, where the value for MAE decreases when increasing the chain length. This reduction in MAE can be understood in terms of the addition of Mn atoms that tend to align their magnetic axis perpendicular to the Fe one. These results qualitatively agree with the results of anisotropic spin Hamiltonians where the total magnetic anisotropy is approximated by adding the individual contributions of $D$ and $E$. This approximation does not capture the changes in geometry of the chain with increasing length. Nevertheless, adding up the MAE contributions of each atom leads to overestimations of MAE that are below the accuracy of our DFT calculations.

The study of the magnetic anisotropy in these chains reveals their composite structure. All FeMn$_x$ for odd-$x$ show a quasipin of 1/2, which should present zero anisotropy if these chains were macrospins of spin 1/2, see Eq. (2). Our DFT calculations show sizable anisotropies that underscore the complexity of the magnetic states of these antiferromagnetic structures.

E. Magnetic excitation energies

The magnetic structure of the chains can be obtained by studying the inelastic electron tunneling spectra (IETS) [9] obtained with the STM. Figure 6 shows the differential conductance obtained for the different FeMn$_x$ chains. The features that appear in these spectra are due to magnetic excitations, very similarly to the ones of Mn$_x$ chains shown in Ref. [22]. Nevertheless, there are noticeable differences regarding both the peak at zero bias for the odd-$x$ chains and the detailed structure of the steps. Indeed, even-$x$ chains are singlets in the case of Mn chains, here, however, the ground states of odd-$x$ chains present a doublet ($S \approx \frac{1}{2}$) magnetic structure.

We solve Eq. (1) and study the different magnetic states for each chain using the methodology of Ref. [39]. The study of the ground state is particularly important to understand the behavior of the Kondo physics appearing in Fig. 6 [23]. Indeed, all odd-$x$ chains display a Kondo zero-bias peak, while this peak is absent from the chains with even $x$. Due to the Heisenberg correlations, the experimental ground state is multi-configurational. Solving Eq. (1) with the anisotropy terms, Eq. (2) shows that many of these configurations are $S = \frac{1}{2}$ states, with a weight larger than 20% in the total state. Hence a spin-flip process is possible at zero-energy cost, which explains the appearance of Kondo peaks. Likewise, even-$x$ chains do not have degenerate ground states and Kondo physics is absent. The absence of degenerate ground states for even-$x$ is due to a total spin $S \approx 2$ where the large longitudinal MAE of the Fe atom lifts the degeneracy of the ground state. The presence of anisotropy prevents $S$ from being a good quantum number [23].

The solutions of Eq. (1) permit us to compare the computed data for the chains with the experimental data. Figure 7(a) shows the IETS for the FeMn$_3$ chain measured of the edge Mn atom, and Fig. 7(b) shows two calculations. The first one (blue) performed with the exchange couplings computed from our GGA+$U$ calculations with an effective $U$ value for Mn of 4 eV and for Fe of 1 eV, see Sec. IV C, and adding up the experimental atomic magnetic anisotropies. Unfortunately, DFT seems to underestimate the anisotropy of Fe on Cu$_2$N, see Sec. IV D, therefore we use the experimental values for the anisotropy. The second one (green) scales the exchange couplings by 1.6, using the scaling found for FeMn when the effective $U$ value for Mn was reduced to 3 eV, and keeping the corresponding value of Fe constant. We see that the energy thresholds are in good agreement in the second case, and the solution of Eq. (1) with the third-order perturbation method of Ref. [39] largely reproduces the dynamical phenomenology of the magnetic chain including higher-energy excitations like the one at ±10 meV.

The asymmetry of the main inelastic thresholds found in Fig. 7(a) is treated within the third-order perturbation method by including a potential scattering term in the Kondo scattering [39]. The effect of the potential scattering term is to remove the electron-hole symmetry of the excitation spectra of Eq. (1). In order to fit the experimental spectra, we have used a $J_{\text{Kondo}} \rho = -0.04$ (where $J_{\text{Kondo}}$ is the Kondo exchange coupling with electrons from the substrate and $\rho$ is the density of states at the Fermi energy) and a potential scattering term $U_{\text{Kondo}}/J_{\text{Kondo}} = -0.5$. The results indicate that larger values should be used to reproduce the experimental data, implying the need to go beyond third-order perturbation theory to treat Kondo scattering in FeMn$_3$.

The magnetic behavior of the inelastic thresholds is correctly reproduced by a Zeeman shift. This permits us to extract the value of the gyromagnetic ratio $g$, Fig. 8. For the present case we find that the atomic g’s ($g_{Fe} = 2.1$ and $g_{Mn} = 1.9$ from Ref. [22]) are good approximations to obtain the correct behavior of the magnetic global states with external $B$, Fig. 8. To a large extend, the atomic spin preserves its character, although very entangled due to the sizable Heisenberg exchange interactions.

V. DISCUSSION AND CONCLUDING REMARKS

This work is a detailed experimental and theoretical account of the electronic and magnetic properties of a heterogeneous
type of magnetic atomic chain (FeMn\textsubscript{x} with \(x = 1.6\)) adsorbed on Cu\textsubscript{2}N/Cu (100). The chains are assembled by atom manipulation with an STM tip, and stable configurations are found when the TM atoms (TM = Fe and Mn) sit on top of Cu atoms. Experimentally, it is difficult to assemble chains with \(x > 9\) (we rarely succeeded going beyond \(x = 10\) creating a straight FeMn\textsubscript{x} chain) and theoretically we see that stress builds up as the chain increases size due to the imposed TM-TM distance by the underlying Cu\textsubscript{2}N/Cu (100) substrate. As the chain increases its size, the TM atoms increase their mutual distance and also their distance to the chain, energetically this is translated into a systematic lowering of both the atomization energy and the energy gained by the chain every time a new Mn atom is added.

Upon adsorption, Fe and Mn lose one of their \(\sigma\) electrons in the interaction with the substrate, mainly to form the bond with the neighboring N atoms. There is considerable distortion and hybridization of the d-electron structure but their occupations remain the free-atom ones, leading to magnetic moment values close to the gas phase [13,22].

The experimental data involve the IETS of different chains with detailed information on the excitation energies of the chains. The lower-energy spectral features are due to magnetic excitations of the system as was tested by their magnetic field dependence. In parallel, the values of the Heisenberg Hamiltonian can be obtained from DFT by evaluating the energy of different spin arrangements of the chains. This approach gives us a systematic insight on how the different TM atoms relate to each other in the chain, which are mainly driven by antiferromagnetic superexchange mediated by the nonmagnetic N atoms of the surface. We find that the Fe-Mn couplings are systematically larger than the Mn-Mn ones, and that beyond second neighbors neglect of the magnetic coupling is a very good approximation. Indeed, first neighbors is a sufficient approximation to obtain exchange couplings with an error of 0.2 meV. In the present choice of DFT+U calculations, the computed magnetic exchanges lead to excitation energies smaller than the experimental ones. Unfortunately, there is not a unique way of determining the value of the Hubbard \(U\) for the calculations. Our systematic study of the values show that a change of 1 eV in the value of the Mn \(U\) leads to a \(\sim 60\%\) change in the value of the evaluated couplings yielding good agreement with the experiment. Despite this dependence of the interatomic exchange energies on the value of \(U\), the conclusions of this work do not change for the range of used values. Moreover, the geometrical, stability and magnetic anisotropy results show a negligible dependence with the above values of \(U\).

The TM atoms are subjected to magnetic anisotropies on this surface. Our calculations show that the MAE of the full chain is not just the sum of the MAE’s of each TM atom. Nevertheless, the smallness of Mn MAE renders this approximation acceptable. Despite their different magnetic axis, the very large MAE of the Fe dominates and the Mn

FIG. 7. Inelastic electron tunneling spectra (IETS) of FeMn\textsubscript{3}: (a) experimental and (b) two computed solution using our calculated spin-chain parameters for \((U - J)_{\text{Mn}} = 4\) eV and \((U - J)_{\text{Fe}} = 1\) eV (blue) or scaling the computed exchange coupling by a 1.6 factor as found for FeMn with \((U - J)_{\text{Mn}} = 3\) eV and \((U - J)_{\text{Fe}} = 1\) eV (green), Sec. IV C. The STM tip is placed on the edge Mn atom.

FIG. 8. (a) Inelastic electron tunneling spectra (IETS) for FeMn\textsubscript{3} with a magnetic field applied perpendicular to the surface increasing until 9 T. (b) The behavior of the different magnetic states (black dots) obtained from the experimental figure follows the Zeeman trend expected for a Zeeman term with the atomic \(g\) factors \((g_{\text{Fe}} = 2.1\) and \(g_{\text{Mn}} = 1.9\) from Ref. [22]).
spins orient along the chain following the Fe anisotropy axis. This leads to a collinear solution of the initially noncollinear problem. We have not found any spin canting or frustration although we cannot rule it out for longer chains. Moreover, the study of the magnetic anisotropy in these chains reveals their composite structure. All FeMn$_x$ for odd-$_x$ show a quasi-spin of 1/2. If these chains were spins 1/2, their anisotropy would be strictly zero. However, our calculations show that they have sizable anisotropies in agreement with the complexity of the magnetic states of these antiferromagnetic structures.

This combined experimental and theoretical work gives us direct insight into the different electronic, geometric and magnetic properties of these heterogeneous chains. In particular, we have given an account for the appearance of Kondo peaks and the antiferromagnetic character of these chains, their magnetic anisotropy that permits us to rule out a macrospin behavior, as well as the accumulated stress that limits the length of the chains.

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