Emergence of half-metallicity in suspended NiO chains

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Contrary to the antiferromagnetic and insulating character of bulk NiO, one-dimensional chains of this material can become half-metallic due to the lower coordination of their atoms. Here we present \textit{ab initio} electronic structure and quantum transport calculations of ideal infinitely long NiO chains and of more realistic short ones suspended between Ni electrodes. While infinite chains are insulating, short suspended chains are half-metallic minority-spin conductors which display very large magnetoresistance and a spin-valve behaviour controlled by a single atom.

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In going from bulk to lower dimensions material properties often change drastically. A recent example is that of interfaces between different insulators which can actually be metallic\cite{1}. Even more recently, it has been predicted theoretically that certain oxygen surfaces of some insulating ceramic oxides can exhibit magnetism and half-metallicity\cite{2}. The ultimate limit in this respect can be found in atomic chains formed in metallic nanocontacts which allow to study the transport properties of one-dimensional systems of atomic size\cite{3}. Due to the lower coordination of the atoms the properties of metallic atomic chains formed in nanocontacts can be remarkably different from those in the bulk. For example, Pt nanocontacts can exhibit magnetic order when atomic chains are formed\cite{4}. More complex one-dimensional systems like carbon-cobalt atomic chains\cite{5} or organometallic benzene-vanadium wires\cite{6} have even been predicted to be half-metallic conductors.

However, not all metals form atomic chains in nanocontacts, although recently, it has been found that the presence of oxygen favours their formation\cite{7}. For example, experiments with Ni nanocontacts\cite{8,9} have never shown evidence of chain formation. Nevertheless, the presence of oxygen in the contact region could possibly lead to the formation of NiO chains. In this context it has also been proposed that the rather moderate magnetoresistive properties of pure Ni nanocontacts could be enhanced considerably by the presence of oxygen adsorbates on the surface of the Ni electrodes\cite{10}. On the other hand, bulk NiO is a common example of a correlated insulator with antiferromagnetic (AF) order (see, e.g., Ref.\cite{11,12}), which remains insulating even above the Néel temperature when the AF order is lost. Thus it is not at all obvious whether or not oxidized Ni nanocontacts or NiO chains should be conductors.

In this Letter we investigate the electronic and magnetic structure and the transport properties of one-dimensional NiO chains, both idealized infinite ones and more realistic short ones suspended between Ni nanocontacts. Anticipating our most important results our \textit{ab initio} quantum transport calculations show that short NiO chains suspended between Ni nanocontacts can become half-metallic conductors, i.e., carry an almost 100\% spin-polarized current. This result holds true even for a single O atom in between Ni electrodes. Consequently, for antiparallel alignment of the electrode magnetizations the transport through the contact is strongly suppressed resulting in very large MR (difference in resistance between antiparallel and parallel alignment of the magnetizations of the electrodes normalized either to the higher resistance value (MR$_1$) or to the lower one (MR$_2$)): MR$_1 \approx 90\%$ and MR$_2 \approx 700\%$, respectively.

The electronic structure of both idealized infinitely long NiO chains and the more realistic short suspended chains have been calculated in the density functional theory (DFT) approximation using the hybrid functional B3LYP\cite{13}, which combines local and non-local exchange. This functional has been successful in describing the electronic and magnetic structure of some strongly correlated materials like bulk NiO. In particular, it describes very well the charge-transfer character, the magnitude of the gap and the magnetic moment of NiO, and predicts the correct AF order\cite{14}. It is important to stress here that due to the insufficient cancellation of the self-interaction in the local exchange functional (DFT) approximation the transport through the contact is strongly suppressed resulting in very large MR (difference in resistance between antiparallel and parallel alignment of the electrode magnetizations normalized either to the higher resistance value (MR$_1$) or to the lower one (MR$_2$)): MR$_1 \approx 90\%$ and MR$_2 \approx 700\%$, respectively.

The electronic structure of infinite NiO chains has been calculated with the CRYSTAL03 program package\cite{17} while for the electronic structure and quantum transport
calculations of the suspended chains we have used our \textit{ab initio} quantum transport package ALACANT \cite{12,19}. The infinite chain calculations were done with elaborate all-electron basis sets for Ni and O \cite{20,21}, similar to those employed for reported HF and B3LYP calculations of bulk NiO \cite{14}, but extended with a diffusive sp-function in the case of Ni and a d-polarization function in the case of O. Using these basis sets we reproduce the B3LYP results for bulk NiO \cite{14}. We have also employed minimal basis sets with effective core pseudo-potentials described in earlier work \cite{14} for both Ni and O, and have found that the results change very little. Thus we have employed these minimal basis sets for the more demanding transport calculations.

The electronic properties of NiO are, to a large extent, determined by the atomic scale properties, like the crystal field splitting of the Ni and O energy levels and the amount of electron charge transferred from Ni to O. The latter, in turn, is determined by the interplay between Madelung binding energy, the ionization potential of Ni, and the electron affinity of O. Due to the lower coordination and the corresponding decrease in Madelung binding energy the electron transfer from Ni to O is less favourable in an atomic chain than in bulk (where the electron transfer is almost complete resulting in an ionic configuration of Ni$^{2+}$O$^{2-}$). The proper starting point to discuss the formation of energy bands in the one-dimensional NiO chain are therefore the univalent ions Ni$^+$ and O$^-$. 

In order to understand how the low coordination affects the atomic properties of the constituting ions we have first performed B3LYP calculations of both a single Ni$^+$ ion and a single O$^-$ ion each in the field of point charges that mimic the crystal field of a one-dimensional chain of univalent Ni and O ions. We find that for both ions the spin-doublet state ($S = 1/2$) minimizes the energy as is the case for the free ions. In Fig. 1 we show schematically the energy levels of Ni$^+$ and O$^-$ in the presence of the point charges for minority spin only. Interestingly, for minority spins, the occupied Ni 3$d_1$ orbitals ($d_{xz}$ and $d_{yz}$) of the Ni$^+$ ion fall energetically in between the occupied and unoccupied O 2$p_1$ orbital ($p_x$ and $p_y$) of the O$^-$ ion. Thus the 3$d_1$ and 2$p_1$ orbitals can form two filled degenerate bonding bands and two degenerate partially filled antibonding bands as indicated in the middle part of Fig. 1. The 3$d_3$ ($d_{xy}$ and $d_{xz-y^2}$) doublet is somewhat above in energy to the 3$d_1$ but cannot hybridize with the oxygen 2$p$ orbitals. The Ni 3$d_0$ ($d_{z^2-r^2}$) and 4$s$ are empty while the O 2$p_0$ ($p_z$) and 2$s$ orbitals are filled and much lower in energy so that no hybridization takes place though symmetry would allow for it.

On the other hand, for majority-spin electrons (not shown) all five Ni 3$d$ orbitals are filled while the 4$s$ is also empty, i.e., the Ni$^+$ valence configuration is 3$d^9$ and not $4s^13d^8$ as for the free Ni$^+$ ion. Moreover, all of the O 2$p$ orbitals are filled so that the Ni$^+$ and O$^-$ ions can only form either completely filled or completely empty bands for the majority spin. Thus the ionic picture suggests that a one-dimensional NiO chain should become a half-metallic conductor where only one kind of spin levels forms conducting bands. In this case only the minority-spin 3$d_1$ bands would conduct so that, according to the proposed classification in Ref. 22, it would be of Type $I_B$. 

\begin{figure}[h]
\centering
\includegraphics[width=\textwidth]{fig1}
\caption{Schematic one-electron energies of a one-dimensional NiO chain in z-direction for minority-spin. To the left and right the orbital energies of an individual Ni$^+$ cation and O$^-$ anion in the crystal field of a one-dimensional Ni$^+$O$^-$ chain are shown. In the center the formation of valence and conduction bands by hybridization of Ni 3$d$ and O 2$p$ orbitals is shown.}
\end{figure}

\begin{figure}[h]
\centering
\includegraphics[width=\textwidth]{fig2}
\caption{(a) Energy per unit cell of infinite NiO chain in dependence of lattice spacing $a$. Black triangles indicate the half-metallic state (HM), grey triangles the molecule-like insulating state with FM order (ML), and black boxes the insulating state with AF order (AF). (b) Band structure of HM state for a lattice spacing of 3.6$\AA$. Solid lines indicate majority-spin bands and dashed lines indicate minority-spin bands. (c) Same as (b) but for ML FM state.}
\end{figure}
Not surprisingly, our calculations for infinite one-dimensional NiO chains (Fig. 2) show that a univalent ionic configuration as an initial guess results in a half-metallic state for large separation of the individual chain atoms (i.e., large lattice spacing of 5Å) as suggested by the ionic picture. However, as can be seen from Fig. 2(a), the half-metallic state is only a metastable state for most values of the lattice spacing. This half-metallic state is “shadowed” by a second state with FM order and insulating character. By successively decreasing the lattice spacing \( a \) of the chain and using the (half-metallic) state of the previous step for the initial guess, the half-metallic state can be generated also for smaller inter-atomic distances which points towards its metastability. Around the equilibrium lattice spacing \( a \approx 3.4\) Å the ground state of the chain has AF order and is of insulating character with a substantial gap of \( \sim 4\) eV like in bulk. When stretched out of equilibrium the FM state and the AF state become comparable in energy until finally, at a lattice spacing of \( \sim 4.2\) Å, the FM state becomes the ground state.

The band structure diagram in Fig. 2(c) shows that the metastable state with FM order corresponds indeed to the half-metallic state suggested by the ionic picture: The half-filled doubly-degenerate conduction band is formed by minority-spin Ni 3d\(_1\) orbitals hybridized with O 2p\(_1\) orbitals while the Ni 3d\(_2\) orbitals do not hybridize with O 2p orbitals and thus form a flat valence band. The lowest-lying empty band is formed by the minority-spin Ni 3d\(_0\) orbital which is slightly hybridized with the Ni 4s orbital. On the other hand the stable state with FM order and insulating character (see band structure in Fig. 2(b)) actually corresponds to the ground state of the NiO molecule which is a \( ^3\Sigma^- \) state. The main difference with the half-metallic state is that now the non-degenerate Ni 3d\(_0\) and 4s orbitals form a minority-spin valence band while the minority-spin doubly-degenerate half-filled antibonding band composed of Ni 3d\(_1\) and O 2p\(_1\) bands is emptied and a substantial gap of \( \sim 3\) eV opens. Thus the infinite chain behaves like an insulator for reasonable values of the chain stretching.

Atomic chains formed in break junctions have a finite length and are suspended between electrodes. It is well known that the contact between the atomic chain and the electrode tip will have considerable effect on the electronic structure of the chain, especially when d-orbitals are involved like is the case here.\(^{10}\) We have thus calculated the electronic structure and transport properties of both a single oxygen atom and a O-Ni-O chain bridging the two tips of a Ni nanocontact as shown in the insets of the right panels of Fig. 3 and Fig. 4. In the case of the single oxygen atom (Fig. 3) the electron transport is almost 100% spin-polarized around the Fermi level for parallel (P) alignment of the magnetizations of the two Ni electrodes. Moreover, an orbital eigenchannel analysis\(^{24}\) reveals that the transport is mainly due to two almost perfectly transmitting minority-spin channels composed of Ni 3d\(_1\) and O 2p\(_1\) orbitals, i.e., they correspond to the doubly-degenerate conduction band of the metastable half-metallic state in the perfect chain. Thus the half-metallic state which was suppressed in the idealized case of the infinite chain emerges in the more realistic situation of a short suspended chain. We can understand this phenomenon in terms of the orbital blocking mechanism proposed earlier in the context of Ni nanocontacts\(^{10,24}\). The highest minority-spin valence band of the insulating state with FM order in the infinitely long chain has a strong contribution from the Ni 3d\(_0\) orbital which is not “compatible” with the geometry of the pyramid shaped Ni contacts, so that this band is blocked and thus cannot be occupied. Instead, the doubly-degenerate band composed of Ni 3d\(_1\) orbitals hybridized with O 2p\(_1\) orbitals is partially filled resulting in the half-metallic state which in the perfect chain is only metastable. Thus the orbital blocking by the contacts actually turns the chain into a half-metallic conductor. Consequently, the conductance is strongly suppressed in the case of antiparallel (AP) alignment of the magnetizations of the Ni electrodes as can be seen from the right panel of Fig. 3 and the MR becomes very large: MR\(_1\) \( \approx 90\)% and MR\(_2\) \( \approx 70\)%.

Varying the distance \( d \) between the Ni tip atoms leads to similar results as those shown in Fig. 3. For the P case the current through the chain is almost 100% spin-polarized with two open minority channels composed of Ni 3d\(_1\) and O 2p\(_1\) orbitals, while for the AP case it is strongly suppressed, resulting in very high MR values between 80% and 90% for MR\(_1\) for \( d \) between 3.0Å and 5.0Å. Geometry relaxations for different values of the tip-tip distance show that for small distances the oxygen atom goes into a zigzag position. The bonding angle decreases with increasing distance until it becomes zero at 3.6Å. Finally, the chain breaks for \( d > 4.8\) Å. Thus the scattering is strong for small distances \( d < 3.6\) Å when the bonding angle is appreciable and for large stretching,
d > 4.2Å, resulting in a considerable reduction in the conduction of the two minority channels.

In longer suspended chains the insulating state with FM order starts to emerge inside the chain and a away from the contacts. As a result the conductance is reduced as can be seen already in the case of the O-Ni-O bridge (Fig. 3). The minority-spin conduction is reduced considerably (~30%) compared to the case of the single oxygen bridge. On the other hand the conduction of the majority-spin channel becomes practically zero (<0.2%). This can be understood by the fact that the residual majority-spin channel conductance of the single oxygen bridge is due to direct hopping of Ni s electrons between the electrodes and therefore vanishes when the distance between the electrodes is large. The finite minority-spin conductance opens up the possibility to an interesting phenomenon: When the middle Ni atom reverses its spin, the conductance drops to nearly zero (see right panel of Fig. 3) since the AF chain is insulating. In other words, this system behaves as a single atom spin valve (by stretching the chain) in suspended chains leads to a strong suppression of the current for AP alignment of the electrodes resulting in very large MR values of MR1 ≈ 90% and MR2 ≈ 700%, respectively. This could perhaps explain to some extend the very large MR values in Ni nanocontacts obtained in some experiments, where oxygen is likely to be present. Finally, the O-Ni-O bridge suspended between Ni electrodes operates as a single atom spin valve where the current flow is controlled by the magnetization of a single atom.

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