Spin-Dependent Quantum Interference Within a Single Magnetic Nanostructure

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Quantum interference is a coherent quantum phenomenon that takes place in confined geometries. Using spin-polarized scanning tunneling microscopy, we found that quantum interference of electrons causes spatial modulation of spin polarization within a single magnetic nanostructure. We observed changes in both the sign and magnitude of the spin polarization between electron waves that are scattered off the boundaries of a nanostructure, forming a standing wave.

When electrons are confined to nanostructures, their dual wave-particle nature comes into full view in experiments. In particular, spatially modulated variations of the electronic local density of states (LDOS) have been observed by scanning tunneling microscopy (STM) (1, 2). These modulation patterns reflect quantum interference between electron waves that are scattered off the boundaries of a nanostructure, forming a standing wave.

In addition to charge, electrons also carry a spin. When confined to magnetic nanostructures, an imbalance between electrons of opposite spin orientations leads to spin-polarized electron waves, giving rise to spin-dependent quantum interference. Recent theoretical studies predicted that local spin polarization on nonmagnetic surfaces as well as on magnetic nanostructures can be spatially modulated (3–5).

We observed spin-dependent quantum interference by means of spin-polarized STM (SP-STM) (6), a technique sensitive to surface magnetization. The nanostructures we study are triangular Co islands located on a nonmagnetic substrate, the (111) surface of copper (4, 7–10). A previous study, which investigated a system similar to ours but under different conditions, indicated the dependence of electron confinement on the spin character of electronic states (9).

The SP-STM measurements were performed at a temperature of 8 K. To detect magnetic contrast, we used W tips covered with magnetic materials (Cr-Co-W tips) and external magnetic fields up to \( B = 4 \) T. To explore the spatial distribution of the spin polarization, we measure the spatially resolved map of the differential tunneling conductance (dI/dV) asymmetry, \( A_{dI/dV} \), defined by

\[
A_{dI/dV} = \frac{dI/dV_{AP} - dI/dV_{P}}{dI/dV_{AP} + dI/dV_{P}}
\]

where \( dI/dV_{AP} \) and \( dI/dV_{P} \) are the dI/dV signals measured with the tip and sample magnetization in antiparallel (AP) and parallel (P) configurations, respectively. The dI/dV signal depends on the relative orientation of the tip and sample magnetizations, which is described in the generalized Tersoff-Hamann model (11, 12). Using that model, we can link the dI/dV asymmetry \( A_{dI/dV} \) to the spin polarization of the tip, \( P_T \), and of the sample at the tip apex position, \( P_S \) (10, 13):
Fig. 2. (A and B) Two \(dI/dV\) images of the Co island in Fig. 1, the basis for the \(dI/dV\) asymmetry map in (C). Both images were recorded at \(B = -1.1\ T\), but with different magnetization configurations between the magnetic tunneling tip and the Co island: antiparallel (A) and parallel (B). The insets represent the antiparallel (AP) and parallel (P) configurations. \(V = +0.03\ V\), \(V_{stab} = +0.5\ V\), and \(I = 1.0\ nA\), where \(V\) is the bias voltage at which the \(dI/dV\) signal is recorded and \(V_{stab}\) is the bias voltage to stabilize the tip before the feedback loop is opened (10). (C) \(dI/dV\) asymmetry map calculated using Eq. 1 from the images in (A) and (B).

We used Eq. 1 to calculate the \(dI/dV\) asymmetry map (Fig. 2C) from the two \(dI/dV\) images. The \(dI/dV\) asymmetry near the Fermi level at \(V = +0.03\ V\) is strongly position-dependent within the Co island. At the edge of the island it is negative, whereas the inner part of the island shows largely positive values. This result can be explained by the existence of a rim state, which is localized spatially around the edges of the Co island and energetically around the Fermi level (9). The rim state originates from a minority d state (9), whereas the inner part of the island mainly has the opposite spin character around the Fermi level and is of majority s-p surface state (4, 7). The magnetization of the tip did not change direction, and the applied bias voltage was fixed at \(V = +0.03\ V\) during the two measurements of the \(dI/dV\) image. Thus, we can assume that the spin polarization of the tip, \(P_{t}\), is constant and the \(dI/dV\) asymmetry is proportional to the spin polarization of the sample—that is, \(A_{dI/dV} \propto P_{s}\). Consequently, we conclude that the inner part of the Co island exhibits a positive \(dI/dV\) asymmetry because the electronic state with the minority spin character is dominant, and that the edge of the Co island exhibits a negative \(dI/dV\) asymmetry because the electronic state with the minority spin character is dominant. To interpret the spatial modulation in the \(dI/dV\) asymmetry (Fig. 2C), which is not due to the atomic structure of the Co island, we focus our discussion on the inner part of the island and analyze the system theoretically.

Our analysis uses density functional theory implemented in the multiple-scattering Korringa-Kohn-Rostoker Green’s function method (10, 20) to calculate spatially resolved maps of the LDOS above a triangular two-atomic-layer Co island on Cu(111) for the majority (Fig. 3A) and minority (Fig. 3B) spins at the Fermi level. The LDOSs for the majority spins are spatially modulated inside the triangular Co island. We ascribe this to a standing wave induced by the electronic confinement of the free electron–like surface state. Minority spins have a constant LDOS, consistent with the understanding that they are mainly due to energetically and spatially localized d states, which are hardly influenced by the electronic confinement (4). We extract the spatial distribution of the spin polarization above the Co island, \(P_{Co}\) (Fig. 3C), from the two calculated LDOS maps as

\[
P_{Co} = \frac{n_{↑} - n_{↓}}{n_{↑} + n_{↓}}
\]

where \(n_{↑}\) and \(n_{↓}\) are the LDOS in the vacuum region above the Co island for the majority and minority spins, respectively (13). To identify the origin of the largely positive spin polarization observed in Fig. 3C, we present two line profiles (Fig. 3D) of the LDOS maps of Fig. 3, A and B. At the Fermi level, the spatially modulated LDOS of the majority spin is larger everywhere in the inner part of the island than the spatially flat LDOS of the minority spin. This leads to a positive spin polarization, \(P_{Co} > 0\) in Eq. 3, in this region.

Comparing Figs. 2C and 3C, we find good agreement between the measured \(dI/dV\) asymmetry map and the calculated spin polarization map within the inner part of the Co island. Therefore, the \(dI/dV\) asymmetry map of the Co island can be interpreted as follows: (i) The \(dI/dV\) asymmetry map qualitatively shows a spatial distribution of the spin polarization on the Co island at a certain energy (Figs. S2 and S3) (21). (ii) The modulation pattern observed in the \(dI/dV\) asymmetry map mainly originates from that in the LDOS of the majority spin, which is ascribed to the electron quantum confinement of the free electron–like s-p surface state. Thus, the spin polarization within the Co island is spatially modulated because of spin-dependent quantum interference.

Next, we examine the energy dependence of the \(dI/dV\) asymmetry maps of the same Co island to ensure the interpretation (ii) described above. The modulation pattern in the \(dI/dV\) asymmetry maps should change with electron energy. The \(dI/dV\) asymmetry maps (Fig. 4, B to E) show clear spatial modulations, similar to those in the corresponding \(dI/dV\) images (fig. S4), with modulation patterns strongly dependent on energy. This is easily understood by considering the origin of the modulation pattern, which is electron confinement of the free electron–like s-p surface state. The s-p surface state starts around \(-0.2\ eV\) below the Fermi level and exhibits a parabolic dispersion with a positive effective mass (7, 9). With increasing energy, the parallel wave vector \(k_{z}\) of the s-p surface state increases; that is, the wavelength \(\lambda\) of the standing waves becomes shorter. The modulation pattern observed in the \(dI/dV\) asymmetry maps shows the same trend as the standing wave pattern in the \(dI/dV\) images. This result corroborates the interpretation (ii) of the \(dI/dV\) asymmetry map.
asymmetry maps. Unfortunately, we also find in Fig. 4, B to E, that the sign of the $dI/dV$ asymmetry changes with respect to energy, whereas the map at $V = +0.25$ V (Fig. 4D) reveals oscillatory change of the sign as a function of position.

To explore the physics behind these surprising results, we plotted a calculated spin-resolved LDOS above a bilayer Co film on Cu(111) as a function of energy (Fig. 4A). Here we focus on the energy range where the free electron–like s-p surface state arises. The LDOS for the majority spin monotonically increases from $E = -0.25$ eV with increasing energy, which is ascribed to the free electron–like s-p surface state. In contrast, the LDOS for the minority spin shows two energetically localized d states around $E = -0.25$ eV and $+0.30$ eV, and it is featureless in the interval between them. The spin-resolved LDOS plots reveal that the dominant spin character strongly depends on energy. We can never conclude that the minority spin character is dominant, even in the energy range where the standing wave pattern, which results from the majority s-p surface state, is clearly observed. We must precisely take into account the dominant spin character at a given energy as revealed by the calculations to evaluate the spatial distribution of the $dI/dV$ asymmetry within the Co island.

At $V = -0.15$ V (Fig. 4B), the inner part of the Co island has a modulated negative value of the $dI/dV$ asymmetry. The spin-resolved LDOS plot (Fig. 4A) shows that the minority spin state is dominant at the corresponding energy, $E = -0.15$ eV, as the localized minority d state exists. Thus, the spatially flat LDOS of the minority spin exceeds the spatially modulated LDOS of the majority spin inside the Co island. This leads to the negative spin polarization inside the island, $P_{Co} < 0$, which comes from a negative numerator in Eq. 3 (i.e., $n_1 - n_1 < 0$), in agreement with the calculated spin polarization map (Fig. 4F). We note that the periodicity of the standing waves in the energy range considered, $\pm 1.5$ nm, is much larger than the distance between neighboring Co atoms, $\pm 0.2$ nm.

At $E = +0.25$ eV, where the spin-resolved LDOS plots of the majority and the minority spins cross, we expect that the net spin polarization should be zero within the Co island. The spatial distribution of the LDOS of the majority spin is, however, not flat but modulated. The LDOS of the majority spins is larger than that of the minority at a convexity of the standing wave, and the resulting spin polarization is positive. Correspondingly, the LDOS of the majority spins is smaller than that of the minority at a concavity of the standing wave, and the resulting spin polarization is negative. Where the LDOS of the majority spins is comparable to that of the minority at a node of the standing wave, the resulting spin polarization is zero. Therefore, the spin polarization map shows a spatial modulation and the sign of the spin polarization oscillates as a function of position, depending on the magnitude of the modulated LDOS of the majority spin (Fig. 4H).

We add a color map to Fig. 4A to clarify the energy window where the experimental $dI/dV$ asymmetry map exhibits only positive, only negative, or both signs of the $dI/dV$ asymmetry within the inner part of the Co island. We find that the dominant spin character at a given energy governs the sign of the $dI/dV$ asymmetry and its spatial distribution.

The finding that not only the magnitude but also the sign of the spin polarization is spatially and energetically modulated within a single magnetic nanostructure implies that we can easily change or control local spin polarization on a magnetic nanostructure by changing the energy at a given position, or the position at a given energy.

Our method for extracting the $dI/dV$ asymmetry map, which requires that the relative magnetization directions of sample and magnetic tip are experimentally determined by in-field STM, allows us to qualitatively visualize the spin polarization of a single nanostructure on an angstrom scale. Recent ab initio calculations predicted that the spin polarization of surface-state electrons on Cu(111) caused by magnetic adatoms can be enhanced within a Cu corral, and the quantum confinement of surface electrons within corrals or islands can be exploited to tailor the exchange interaction between magnetic adatoms (3, 22). When combined with the capability of the STM to manipulate adatoms and assemble engineered nanostructures (23, 24), our method offers a way
Multiple Functional Groups of Varying Ratios in Metal-Organic Frameworks

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We show that metal-organic frameworks (MOFs) can incorporate a large number of different functionalities on linking groups in a way that mixes the linker, rather than forming separate domains. We made complex MOFs from 1,4-benzenedicarboxylate (denoted by “A” in this work) and its derivatives -NH2, -Br, -(Cl)2, -NO2, -(CH3)2, -C6H4, -(OC6H4)2, and -(OC3H5)2 (denoted by “B” to “I,” respectively) to synthesize 18 multivariate (MTV) MOF-5 type structures that contain up to eight distinct functionalities in one phase. The backbone (zinc oxide and phenylene units) of these structures is ordered, but the distribution of functional groups is disordered. The complex arrangements of several functional groups within the pores can lead to properties that are not simply linear sums of those of the pure components. For example, a member of this series, MTV-MOF-5-EHI, exhibits up to 400% better selectivity for carbon dioxide over carbon monoxide compared with its best same-link counterparts.

Our strategy to making MTV-MOFs is to assemble their structures from links with different functional groups whose orientation, number, relative position, and ratio along the backbone (metal-oxide and phenylene units) can be controlled by virtue of the unchanged length of the links and its unaltered connectivity (Scheme 1). Such a construct can be viewed as having a primary structure composed of the simple repeating pattern of metal-oxide joints and organic links and a “complex” secondary structure formed by multivaried arrangements of many functional groups that are covalently bound to the links. In this way, each of the pores within the MOF would have an array of functionalities pointing into its center. Accordingly, the sequence of such functionalities and the frequency with which certain ones appear in the sequence will endow the pores with a new level of complexity that far exceeds any held by that of the original same-link MOFs—an aspect that may allow fine-tuning of the pore environment with favorable implications on properties.

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References and Notes
10. See supporting material on Science Online.
13. We note that the LDOS in vacuum strongly depends on the distance between the STM tip and the scanned surface (25).
21. We note that the spin polarization shown here is not identical with the local magnetic moment of the sample because it is obtained at a certain energy.

References

See supporting material on Science Online.

Our present work (as well as the work of others) has shown that MOFs with two mixed links can be prepared, whereas a recent report showed that four different functionalities can be introduced into one structure by post-synthesis modification (2–7). These approaches are either confined to only two links or severely limited by having complete reactions at the links: multiple variations in link ratios and functionalities in these systems were not demonstrated. The present report describes a general method for producing crystalline MOF materials that combine sets of two to eight links of different functional groups; each set is incorporated into a single structure where the ratio of links is controlled, and the material can be produced with bulk purity. [Hereafter, A, 1,4-benzenedicarboxylate; B, -NH2; C, -Br; D, -(Cl)2; E, -NO2; F, -(CH3)2; G, -C6H4; H, -(OC6H4)2; and I, -(OC3H5)2.] Specifically, we targeted the cubane MOF-5 structure (8) and combined the acid form of 1,4-benzenedicarboxylate (BDC), NH2-BDC, Br-BDC, (Cl)2-BDC, NO2-BDC, (CH3)2-BDC, C6H4-BDC, (OC6H4)2-BDC, and (OC3H5)2-BDC links (Scheme 2, A to I, respectively) to form the corresponding sets of 18 MTV-MOFs, each having two or more different functionalities [two: MTV-MOF-5-AB, -AC, -AD, -AE, -AF, -AG, -AH, -AI, and -EL; three: MTV-MOF-5-ABC, -AHI, and -EHI; four: MTV-MOF-5-ABCD and -ACEF; five: MTV-MOF-5-ABCH; six: MTV-MOF-5-ABCCH; seven: MTV-MOF-5-ABCEGHI; eight: MTV-MOF-5-ABCEFEGHI (Scheme 2)]. We describe their isolation as single phases, the structure of their MOF backbone, and their porosity, and show that this multivaried link synthetic strategy is useful for introducing functionalities [such as NO2-BDC and (Cl)2-BDC] into the MOF-5 type structure (MTV-MOF-5-AD and -AE) that do not form this structure when used alone. We also report our initial findings that members of this series (MTV-MOF-5-AHI and -EHI) show that the “whole is better than the sum of its parts,” as evidenced by the fourfold enhancement of gas adsorption and separation properties of the multivaried

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References
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