Impurity states and interlayer tunneling in high temperature superconductors

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We argue that the Scanning Tunneling Microscope (STM) images of resonant states generated by doping Zn or Ni impurities into Cu-O planes of BSCCO are the result of quantum interference of the impurity signal coming from several distinct paths. The impurity image seen on the surface is greatly affected by interlayer tunneling matrix elements. We find that the optimal tunneling path between the STM tip and the metal (Cu, Zn, or Ni) \(d_{x^2-y^2}\) orbitals in the Cu-O plane involves intermediate excited states. This tunneling path leads to the four-fold nonlocal filter of the impurity state in Cu-O plane that explains the experimental impurity spectra. Applications of the tunneling filter to the Cu vacancy defects and “direct” tunneling into Cu-O planes are also discussed.

Recently J.C. Davis and collaborators applied the STM technique to image single Zn and Ni impurities in optimally doped BSCCO [1,2]. These experiments proved that one can image single impurity states in an unconventional superconductor and demonstrated the highly anisotropic structure of these states.

Although it appears that on a gross scale these findings can be understood in terms of a conventional \(d\)-wave superconductor perturbed by potential scattering, upon closer inspection problems of principle seem to arise. The impurity states observed by STM are characterized by two main features: (1) energy and width of the impurity-induced resonance in the density of states (DOS), and (2) the spatial structure of the resonance. While the DOS seems to be satisfactorily described by a single-site impurity model [3], the real space distribution of intensity cannot be fit by this model. The main problem with the Zn impurity image seen in the STM experiments is that the intensity of the signal on the impurity site is very bright, which is at odds with the unitary scattering off Zn. We remind that Zn\(^{2+}\) has a closed \(d\) shell and hence, exactly on the impurity site the scattering potential is very strong. Unitary scattering is equivalent to the hard wall condition for the conduction states and therefore no or very little intensity of electron states is expected on the Zn site. A similar problem arises also with explaining the Ni-induced resonance.

Here we demonstrate that these problems find a natural resolution in terms of the specific way in which the local density of states of the cuprate planes is probed in the STM experiments. We argue that the quantum-mechanical nature of the tunneling from the STM tip into the Cu-O layer that hosts impurity requires tunneling through the uppermost insulating Bi-O layer which effectively filters the signal. Surprisingly, similar filtering should also take place even in the case of “direct” tunneling into Cu-O plane. Such non-local tunneling has profound consequences for the real space image of the impurity state seen by STM.

There are two major types of the tunneling routes between the STM tip and the conducting orbitals in the Cu-O plane: (a) \textit{Direct} tunneling due to the overlap between the tip and the planar \(3d_{x^2-y^2}\) wavefunctions, and (b) \textit{indirect} tunneling through intermediate excited (occupied or empty) states, Fig. 1a. The direct tunneling probability over the experimentally relevant distances of about 10Å is however exponentially small [3]. On the other hand, the importance of the excited states for mediating STM tunneling follows directly from the Bi-O topographs that clearly show the positions of the nominally insulating Bi atoms on the surface of BSCCO. The analysis of the topographs implies that the excited Bi orbitals \textit{focus} the flow of the tunneling electrons. Therefore, we argue that the indirect tunneling via overlapping intermediate orbitals is the dominant tunneling mechanism.

The strongest indirect tunneling channel involves intermediate states that have the largest overlaps. In the case of BSCCO, these are the orbitals that extend out of the planes, such as \(4s\) or \(3d_{x^2-y^2}\) of Cu and \(6p\) of Bi. Furthermore, only the states with zero in-plane orbital momentum have non-zero overlap with the apical oxygen \(2p_z\) and \(3s\) orbitals that play important role in the interlayer communication. Being radially symmetric in the Cu-O plane, such states \textit{cannot} couple to the relevant \(3d_{x^2-y^2}\) states \textit{on-site}. They do couple however to the \textit{neighboring} \(3d_{x^2-y^2}\) orbitals through the \(d\)-wave-like \textit{fork} (Fig. 1b). The resulting tunneling amplitude is

\[
M_{i,j} \sim \Psi_{i+1,j} + \Psi_{i-1,j} - \Psi_{i,j+1} - \Psi_{i,j-1},
\]

where \(\Psi_{i,j}\) is the impurity state wavefunction on site \((i,j)\). Hence, the symmetry analysis leads to the surprising conclusion: Tunneling on top of a particular Cu (or Zn or Ni) atom in the Cu-O plane \textit{does not} probe its \(3d_{x^2-y^2}\) orbitals, but rather measures a linear combination of its neighbors. The tunneling matrix element determines the intensity of the impurity signal \(F[\Psi_{i,j}] = |M_{i,j}|^2\). Calculating the spectral intensities in the vicinity of the impurity using this filtering function
reproduces the features seen in the experiments in great detail (Figs. 3 and 1). The aforementioned anomalies find an explanation in terms of interferences associated with the non-local way in which the electronic states are probed. The described filtering mechanism is related to the one responsible for the interlayer tunneling in the bulk cuprates \cite{Hwang2002}. An alternative non-local filter based on the incoherent direct tunneling from the tip into Cu-O plane was recently studied by Zhu et al. \cite{Zhu2002}. The angular effect of the filter on the far asymptotic of the impurity states was discussed earlier \cite{Fe autofocus 1999}.

![Diagram](image-url)

**FIG. 1.** (a) Tunneling from STM tip into $3d_{x^2-y^2}$ orbitals in the Cu-O layer occurs through the virtual energy states of Bi and Cu. The tunneling processes involve both empty (top) and occupied (bottom) intermediate states. All such processes contribute coherently to the tunneling amplitude from the tip to the $3d_{x^2-y^2}$ orbitals. (b) Tunneling through a particular Bi atom does not probe the $3d_{x^2-y^2}$ orbital of the metal atom (Cu, Zn or Ni) right underneath it due to the vanishing overlap. Instead, the tunneling involves orbitals that extend out of the planes, such as $4s$ (shown) or $3d_{xz,z^2-r^2}$ of Cu and $6p_z$ of Bi. These orbitals are symmetric in the Cu-O plane and hence couple to the neighboring metal $3d_{x^2-y^2}$ orbitals through the $d$-wave-like fork, Eq. (6).

Similar to the Zn case, the Ni impurity tunneling intensity is maximal at the Bi position immediately above the impurity site. However, there are several important differences between the observed Zn- and Ni-induced states: (1) The energy of the Zn state is close to the chemical potential, $\varepsilon_{Zn} = -2$ mV, while the Ni state energy is larger and is split, $\varepsilon_{Ni} = 9$ and 18 mV, (2) the Zn state appears only on the negative bias, while the Ni state shows up both on positive bias and the symmetric negative bias. In this letter we demonstrate that these experimental features can also be reproduced within the standard theory of the impurity states \cite{Fe autofocus 1999}, with the spatial structure of the states being reproduced by properly taking into account the fork effect.

The starting point of our model is a two-dimensional mean-field (MF) Hamiltonian with the nearest neighbor attraction, $V$, which yields $d$-wave superconductivity in the range of dopings close to half-filling.

$$H_0 = - \sum_{i,j,\sigma} t_{ij} c_i^{\dagger}_\sigma c_j^\sigma + \sum_{\langle ij \rangle} c_i^{\dagger} c_j^\dagger \Delta_{ij}^* + \text{h.c.} \quad (2)$$

Here, $\Delta_{ij} = V(c_i^{\dagger} c_j^\dagger)$ is the self-consistent MF superconducting order parameter. The hopping $t_{ij}$ equals $t$ for nearest neighbors and $t'$ for for the second-nearest neighbor sites $i$ and $j$. The parameters that are relevant for BSCCO are $t = 400$ meV, $t' = -0.3t$. To match the amplitude of the superconducting gap in optimally-doped BSCCO, which is about 40 meV, we choose the attraction $V = -0.525t$. The chemical potential is chosen to yield 16% doping ($\mu = -t$).

The local impurity is introduced into Hamiltonian Eq. (2) by modifying the electron energy on a particular site \cite{Fe autofocus 1999}. The corresponding addition to the Hamiltonian is

$$H_{\text{imp}} = V_{\text{imp}}(n_{0\uparrow} + n_{0\downarrow}) + S_{\text{imp}}(n_{0\uparrow} - n_{0\downarrow}) \quad (3)$$

The first term is the potential part of the impurity energy that couples to the total electronic density on site 0, and the second term describes the magnetic interaction of the impurity spin and the electronic spin density on the same site. We assume that the impurity spin is large and can be treated classically, as if it were a local magnetic field. The goal is to determine $V_{\text{imp}}$ and $S_{\text{imp}}$ so as to match both the location of the impurity states within the gap and the spatial distribution of their intensity.

First let us analyze the position of the impurity energy level as a function of the impurity potential. We solve the MF equations self-consistently on a square lattice with periodic boundary conditions. The results are presented in Figure 2. All impurities placed in a superconductor generate quasiparticle weight both on positive and symmetric negative biases. The sign of the energy level is defined based on where the majority of the quasiparticle weight resides. Attractive impurities produce energy levels lying below the Fermi surface, while repulsive ones generate states with positive energy. In the limit of a very strong impurity potential both attractive and repulsive impurities generate identical states with a small
residual energy related to the amount of the particle-hole symmetry breaking, caused by the specifics of the band structure and doping [12].

FIG. 2. Impurity energy level position as a function of the potential impurity strength, \( V_{\text{imp}} \). The blue (top) line corresponds to repulsive impurities, which produce positive energy levels (\( \varepsilon_{\text{imp}} > 0 \)); the red line is for attractive impurities, which generate negative energy states (\( \varepsilon_{\text{imp}} < 0 \)). For strong impurity potential both energies converge to the same small value determined by the amount of particle-hole symmetry breaking.

The analysis of the energy level positions implies that the Zn impurity can be associated with a strong attractive potential, \( V_{\text{Zn}} = -11t = -4.4 \text{ eV} \) and \( \varepsilon_{\text{Zn}} = -0.005t = -2 \text{ meV} \). Ignoring for now the level splitting, the Ni case can be associated with a relatively weak repulsive impurity, \( V_{\text{Ni}} = t \) and \( \varepsilon_{\text{Ni}} = 0.0443t = 18 \text{ meV} \). These impurity strengths are in agreement with the general band structure arguments. Zn\(^{2+}\) ion has 10 electrons that completely fill \( d \) orbitals. Hence, the \( d_{x^2-y^2} \) orbital of Zn, relevant for interaction with Cu-O plane orbitals, is deep below the Fermi surface. On the other hand, Ni\(^{2+}\) has 8 electrons in the \( d \) shell, with the \( d_{x^2-y^2} \) being unoccupied, but with a small energy, given by the level splitting within the \( d \) shell.

The spatial distribution of the spectral intensity corresponding to the Zn impurity is shown in Figure 3. The top two plots show the intensity \( A_{\text{CuO}}^{i,j} \) as it would be seen if STM tip were directly imaging DOS in the Cu-O plane. The intensity is related to impurity state wavefunction \( \Psi \) on site \((i,j)\),

\[
A_{i,j}^{\text{CuO}} = |\Psi_{i,j}|^2. \quad (4)
\]

The intensity on the impurity site is suppressed due to the strong impurity potential. The bottom two plots correspond to imaging through the top Bi-O layer. They are obtained by applying a filtering function \( F[\Psi] = |M_{i,j}|^2 \) to the impurity state wavefunction \( \Psi \). The effect of the filtering function is to produce the intensity

\[
A_{i,j}^{\text{BiO}} \propto |\Psi_{i+1,j} + \Psi_{i-1,j} - \Psi_{i,j+1} - \Psi_{i,j-1}|^2. \quad (5)
\]

Indeed, the intensity on the Bi-O layer is maximized on the impurity site due to the interference of the contributions from the impurity’s nearest neighbor sites.

The structure of the Ni-induced state is more complicated than the Zn case. It appears on both positive and negative biases. In addition, there is a peak splitting on each bias. Unlike Zn\(^{2+}\), Ni\(^{2+}\) impurity is magnetic, with spin 1. To simulate the effect of spin we include a non-zero magnetic part of the impurity potential, \( S_{\text{imp}} \neq 0 \), in the Hamiltonian Eq. (3). The spin component introduces level splitting between “up” and “down” spin states. Figure 4a shows the amplitudes of the spin-split states on the impurity sites and its neighbors for \( V_{\text{Ni}} = t \) and \( S_{\text{Ni}} = 0.4t \). The spin-split energy levels are \( \varepsilon_{\uparrow} = 0.052t \) and \( \varepsilon_{\downarrow} = 0.037t \). The total spin-up and spin-down intensities for each bias are shown in Figure 4b. Both figures (a) and (b) correspond to the image affected by the fork filter. The general shape of the states agrees well with the experimental data [3]. The total weight on the negative bias is about factor of 3 smaller than the weight on the positive bias.
FIG. 4. Simulation of Ni as an impurity with weak mixed potential, $V_{Ni} = t$ and $S_{Ni} = 0.4t$. (a) Spectral intensity on the impurity site, its nearest neighbors (nn), and next nearest neighbor (nnn). Red line is spin up, and blue is spin down. (b) Intensity map for the combined spin up and spin down states.

Some aspects of the Ni-impurity state cannot be addressed within the simple framework of our theory. We assumed that the spin of the impurity is static and plays the role of a local magnetic field. Bulk measurements indicate that Ni dopants do not introduce additional magnetic moments. This may be either due to the partial Kondo screening of $S = 1$ Ni spin or due to the local attraction of additional hole (“self-doping”) by Ni$^{2+}$ to form an effective Ni$^{3+}$ complex with spin 1/2. No magnetic signature, combined with the weak potential character of Ni may explain the relatively weak pair-braking effect of Ni doping observed experimentally.

In the case of Zn, we focused on the large local potential scattering off the impurity as a main effect to account for the STM intensity distribution. It is important to mention that Zn substitution also induces local magnetic moment, which is observed in magnetic susceptibility and NMR measurements. The quenching of magnetic moment around Zn can lead to the Kondo resonance near the Fermi surface, as has been proposed by Polkovnikov et al. Below the Kondo temperature, Zn should behave as a unitary scatterer, virtually indistinguishable from either local or extended potential scatterer. We argue however, that the tunneling fork filter should apply regardless of the details of the scattering mechanism.

The tunneling fork mechanism has consequences beyond the settings of the original experiments. Here we point out two of them: (1) the direct tunneling into Cu-O plane, and (2) the structure of the Cu vacancy states. One could assume that the tunneling into the exposed top Cu-O plane should be free of the tunneling fork. However, upon closer inspection it is clear that even in this case, the direct tunneling from tip into the planar $3d_{x^2−y^2}$ orbitals is exponentially weak compared to the indirect tunneling through the s-wave like orbitals of Cu, Zn, or Ni extending out of the plane. Hence, the tunneling fork should apply also in this case, resulting in the same spatial form of the filtered impurity states as obtained above. In the case of Cu vacancy in the Cu-O plane, under assumption that there are no s-wave like orbitals centered at the vacancy site, the tunneling fork mechanism implies that resonance state observed by STM should be the same as for Zn, but with no spectral intensity in the center of the pattern.

In conclusion, we have demonstrated that recently observed Zn and Ni impurity states in BSCCO can be explained by simple model of strong potential impurity in the case of Zn and mixed (potential + spin) impurity in the case of Ni, interacting with a $d$-wave superconducting condensate. The crucial aspect that we have included in the present treatment is the effect of the quantum-mechanical tunneling between the STM tip and the Cu-O planes. The same non-local filter effect should be operable in the case of the “direct” tunneling into Cu-O planes and should have consequences for other types of defects, e.g. Cu vacancies.

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