Doping dependence of gap inhomogeneities at Bi$_2$Sr$_2$CaCu$_2$O$_{8+\delta}$ surfaces

Wei Chen$^{1,4,5}$, Marc Gabay$^2$ and P J Hirschfeld$^3$

1 School of Physics, University of New South Wales, Sydney 2052, Australia
2 Laboratoire de Physique des Solides, Université Paris-Sud 11, Centre d’Orsay, 91405 Orsay Cedex, France
3 Department of Physics, University of Florida, Gainesville, FL 32611, USA
E-mail: w.chen@fkf.mpg.de

New Journal of Physics 14 (2012) 033004 (16pp)
Received 22 October 2011
Published 2 March 2012
Online at http://www.njp.org/
doi:10.1088/1367-2630/14/3/033004

Abstract. We study the inhomogeneity of the pairing gap near the surface of Bi$_2$Sr$_2$CaCu$_2$O$_{8+\delta}$ due to interstitial oxygen dopants. Within the slave boson mean-field theory of a disordered $t$–$t'$–$J$ model, we identify three aspects of the oxygen dopants related to inhomogeneity: (i) the superexchange interaction is locally enhanced in the vicinity of oxygen dopants, which enhances the local pairing gap and reduces the coherence peak; (ii) oxygen dopants donate holes into the CuO$_2$ plane, which reduces the spinon density of states and hence the average gap at large doping; and (iii) holes are locally attracted to the vicinity of oxygen dopants, which causes an impurity bound state and further reduces the coherence peak. The interplay of these mechanisms simultaneously explains the locally enhanced pairing gap around oxygen dopants and the reduction of average gap with increasing oxygen concentration, as observed by scanning tunneling microscopy.

$^4$ Author to whom any correspondence should be addressed.
$^5$ Present address: Max Planck Institute for Solid State Research, Heisenbergstraße 1, 70569 Stuttgart, Germany.
1. Introduction

The effect of disorder in high-$T_c$ cuprates, either intentionally introduced or present as a result of the doping process, has always been crucial for their practical application. For several classes of cuprates, scanning tunneling microscopy (STM) has been particularly useful in clarifying how disorder affects superconductivity [1–6]. In the frequently studied Bi$_2$Sr$_2$CaCu$_2$O$_{8+\delta}$ (BSCCO) system, where disorder is induced primarily by interstitial oxygens which dope the system, STM data show a very pronounced inhomogeneity of superconducting gap over the sample surface. In particular, the gap magnitude is significantly enhanced in the vicinity of oxygen dopants [4].

A phenomenological model was proposed by Nunner et al [7] based on the assumption that the pairing interaction was locally enhanced in the vicinity of oxygen dopants, which successfully captured the feature of locally enhanced superconducting gap and the reduction of the coherence peak in the large gap regions, as well as other statistical correlations. These authors pointed out that such an enhancement could be understood from a strong coupling perspective if the local superexchange $J(r)$ was increased by the presence of a nearby dopant. Maška et al [8] then showed that within a single-band Hubbard model, local variations in site energies as might be due to a dopant perturbation would always lead to an enhanced $J$ within second-order perturbation theory, but similar calculations within a three-band Hubbard model [9] indicated that the sign of $J$ modulation was not universal. Johnston et al [10] found a suppression of the superexchange in Cu–O cluster calculations with point charge estimates of Madelung energies, but pointed out that enhancement could occur if the electron–phonon coupling to the $c$-axis oxygen modes were accounted for. From a weak-coupling perspective, Foyevtsova et al [11] considered the change in local electronic structure due to an interstitial O in first-principles calculations of BSCCO, and used the results to calculate the spatially averaged spin fluctuation interaction in undoped and doped systems; the enhanced pairing in the doped system was similarly attributed to local enhancement around O interstitials. A somewhat different approach has been pursued by other groups [12, 13], who have discussed the large polarizability of the O interstitial as possibly modulating the overall polarizability of the Cu–O complex and hence the pairing.

Although the inhomogeneous superconducting gap and several important correlations among STM observables are nicely reproduced by the locally enhanced pairing interaction picture at optimal doping, a puzzle remains: if each oxygen dopant locally enhances the superconducting gap, why does the average gap decrease with increasing oxygen concentration? This feature has been quantified in [5], where empirically not only is the average gap found to decrease with oxygen concentration, but also the spatial distribution of the superconducting gap...
is observed to be narrower at large doping. This apparent discrepancy indicates that additional ingredients are necessary in order to understand the doping dependence of the inhomogeneous superconducting gap. A natural assumption is that at least one of these additional ingredients must be global electronic correlations, which affect the average superconducting gap over a rather large scale, but still allow the local gap to be enhanced around each oxygen dopant.

In this work, we use slave boson mean-field theory to study the inhomogeneous pairing gap on the surface of BSCCO, and show that besides the locally enhanced pairing interaction, the holes donated by oxygen dopants have a dramatic effect on the pairing gap. We find that at large oxygen concentration, the hole doping level increases accordingly, which reduces the density of states (DOS) of spinons and causes the average gap to drop. The advantage of slave boson mean-field theory is that it treats disorder and strong correlations on an equal footing, since the inclusion of strong correlations is necessary to explain the reduction of DOS. A theory of this kind has been applied to studying YBa$_2$Cu$_3$O$_{6+x}$ (YBCO) in the presence of a point-like repulsive impurity \cite{14}, where the repulsion of holes and the reduction of the fermionic bandwidth in the vicinity of the impurity influence the charge and spin responses significantly, yielding a nuclear magnetic resonance (NMR) spectrum consistent with experiments. In essence, oxygen dopants in BSCCO serve as impurities of the opposite kind: their Coulomb potential attracts holes which, in the slave boson description, promotes the enhancement of the fermionic bandwidth. We will demonstrate that the enhancement of fermionic bandwidth serves as another mechanism to reduce the coherence peak in the large gap regions.

A similar method \cite{15} has been proposed previously to study the effect of oxygen dopants in BSCCO, where screening of the ionic Coulomb potential is studied in detail, and is argued to be the origin of electronic inhomogeneity. However, without a locally enhanced pairing interaction, the superconducting gap is found to be anticorrelated with oxygen dopants, in contrast to the observation of STM. It was also proposed that postulating two types of oxygen dopants, one playing the role of the Coulomb defect and the other purely controlling the doping, can resolve the O-gap anticorrelation issue \cite{16}. However, based on various microscopic calculations \cite{8–13} that support the possible enhancement of the local pairing interaction by oxygen dopants, we believe that it is important to include this effect in a phenomenological way, and treat each oxygen dopant as both a Coulomb center and a source of charge. We will show that the positive correlation between pairing gap and oxygen dopants is recovered by properly accounting for various energy and length scales in the problem. It is also proposed that slave boson treatment naturally enhances the pairing interaction around oxygen dopants, without any additional microscopic mechanisms such as spatially varying atomic level \cite{17}. The argument is based on the renormalization of hopping integral $t_{ij}$ and its relation to the superexchange interaction $J_{ij} = 4t_{ij}^2/U$. However, this mechanism gives a very small enhancement of $J_{ij}$, since we found that the hopping integral is only renormalized by about 20% near oxygen dopants, which cannot account for the large gap inhomogeneity observed in STM. The inclusion of a phenomenological, locally enhanced pairing interaction in the framework is therefore still necessary for the development of the theory. It is also worth mentioning that BSCCO exhibits a unique incommensurate structural modulation. However, the position of the oxygen dopants identified by STM, as well as the pairing enhancement near oxygen dopants, does not seem to have a direct correlation with the incommensurate modulation \cite{4}. We assume that this modulation does not play an important role in pairing inhomogeneity and can be safely ignored in the calculation.
The structure of this paper is as follows. In section 2 we revisit the enhanced pairing interaction picture in [7] and study the change of local bandwidth due to strong correlations phenomenologically. This section essentially illustrates all the physical effects that occur in the vicinity of each oxygen dopant. In section 3, we introduce the slave boson mean-field treatment for homogeneous cuprates and demonstrate the well-known reduction of average gap due to hole doping. In section 4, oxygen dopants are introduced in the slave boson treatment to study disorder and correlations on an equal footing. Section 5 presents the conclusions.

2. Phenomenological model of a single oxygen dopant

In this section, we follow the strategy of [7] to study a single oxygen dopant within a Hartree–Fock–Gor'kov mean-field approach. The motivation is to clarify the influence of oxygen dopants on the local density of states (LDOS), in the simplest non-self-consistent manner. In comparison with the fully self-consistent treatment in section 4, only the fermionic degrees of freedom are considered in this section, since they are essential for local pairing, and the effect of strong correlations is discussed phenomenologically. We consider the following impurity Hamiltonian:

\[
H = \sum_{ij\sigma} (-t_{ij} - d_{t_{ij}}) c^\dagger_{i\sigma} c_{j\sigma} + \sum_{i\sigma} (V_i - \mu) n_{i\sigma} + \sum_{\langle ij \rangle} (\Delta_{ij} + d\Delta_{ij}) c^\dagger_{i\sigma} c_{j\sigma} + \text{h.c.} \tag{1}
\]

For simplicity and consistency with [7, 18], in this section we assume that the oxygen dopant is located on top of a random Cu site (this is not quite correct [19]), and we choose \( t = 1 = 400 \text{ meV} \) for nearest-neighbor and \( t' = -0.3 \) for the next-nearest-neighbor hopping, and \( \mu = -1.0 \). The homogeneous pairing amplitude is \( \Delta_{ij} = \Delta_0 (\delta_{i=r, j=x} - \delta_{i=r, j=y}) \), with \( \Delta_0 = 0.1 \). In [7], it is also shown that if only the electronic (fermionic) degrees of freedom are accounted for, the usual association of an on-site Coulomb potential \( V_i \) with each oxygen dopant gives the form of the electronic structure, including a highly inhomogeneous low-energy LDOS, that is inconsistent with STM experiments. We therefore neglect the on-site part of the impurity potential, \( V_i = 0 \). Here we focus on the effect of hopping impurities \( d_{t_{ij}} \) and the off-diagonal (pairing channel) impurities \( d\Delta_{ij} \), and assume that they only differ from the bulk value on the four bonds connected to impurity site \( r \), \( d_{t_{ij}} = d_{t_{ji}} = d t (\delta_{i=r, j=x} + \delta_{i=r, j=y}) \) and \( d\Delta_{ij} = d\Delta_{ji} = d\Delta (\delta_{i=r, j=x} - \delta_{i=r, j=y}) \).

For comparison with [7, 18], we first perform simulations for a single off-diagonal impurity. As shown in figure 1(a), one clearly sees an enlarged gap and reduced coherence peak for the case of local pairing enhancement \( d\Delta / \Delta_0 = 1 \). The off-diagonal impurity case obviously captures the enhancement of the gap by oxygen dopants. We then perform simulations for the hopping impurity, for both hopping enhancement and reduction cases. The motivation comes from the slave boson treatment of the BSCCO surface, where the spinon hopping amplitude is enhanced near oxygen dopants due to the accumulation of holons, as will be addressed in section 4. Here we treat this effect at the phenomenological level by using a hopping amplitude locally modulated by roughly \( dt / t \sim 10\% \) as found by the microscopic theory discussed in section 4. Figure 1(b) shows the simulation for both hopping enhancement and reduction on four bonds connected to the oxygen dopant. The LDOS on the impurity site clearly indicates that hopping enhancement reduces the coherence peak and vice versa, while the gap size remains unchanged. The LDOS calculated over a wider energy range, as shown in figure 1(b), demonstrates that the reduced coherence peak is due to additional bound states formed on the
Figure 1. LDOS on the impurity site for the phenomenological model described by equation (1), with $t = l = 400\, \text{meV}$, $t' = -0.3$, $V_i = 0$, $\mu = -1$ and $\Delta_0 = 0.1$. Three different impurity models are discussed: (a) pairing enhancement/reduction $d\Delta/\Delta_0 = \pm 1$ on four bonds connected with the impurity site, (b) hopping enhancement/reduction $dt/t = \pm 0.1$ on four bonds and (c) both pairing and hopping enhancement/reduction. Insets in (a)–(c) highlight the LDOS near the chemical potential.

New Journal of Physics 14 (2012) 033004 (http://www.njp.org/)
The simulation that contains both pairing and hopping enhancement is shown in figure 1(c), where one sees that the superconducting gap is enlarged as expected, and the coherence peak is further reduced compared to the case that contains only an off-diagonal impurity. On the other hand, if the impurity locally reduces both the pairing and hopping interactions, then the gap is reduced and the coherence peak height increases. This simulation suggests that the change in local hopping amplitude, whose origin can be traced to the holon accumulation due to strong correlations, plays an important role in the shape of the STM spectrum, particularly the weight of the coherence peak. The calculation that associates a change of hopping amplitude with correlation effects will be demonstrated in section 4.

3. Slave boson mean-field theory in a homogeneous cuprate

Before we introduce the disorder model for BSCCO surface in the next section, we present here the slave boson mean-field model of a homogeneous cuprate. One can regard the phenomenological impurity model in section 2 as a demonstration of how oxygen dopants affect the STM spectrum locally, while the mean-field theory in this section demonstrates how oxygen dopants change the average gap value globally. In the slave boson mean-field treatment at finite doping, spin and charge degrees of freedom can be separated but mutually renormalize each other. The pairing gap is described by the pairing of spinons, which relies on the LDOS of spinon band that in turn is sensitive to hole doping.

We begin with the $t-t'-J$ model treated with the slave boson method

$$
\mathcal{H} = \sum_{ij} -t_{ij} b_i^\sigma f_j^\sigma f_j^\sigma + \sum_{ij} J \left( \tilde{S}_i \cdot \tilde{S}_j - \frac{1}{4} n^f_{i\sigma} n^f_{j\sigma} \right) - \mu_f \sum_{i\sigma} n^f_{i\sigma} - \mu_b \sum_i n^b_i + \sum_i \lambda_i (n^f_{i\sigma} + n^b_i - 1). 
$$

(2)

We choose $t = 400$ meV = 1 and next-nearest-neighbor hopping $t' = -0.4$, a slightly different value from section 2 such that the van Hove singularity is always located close to but lower than the pairing gap. The exchange coupling $J = 72$ meV = 0.18 is smaller than the well-accepted value $J \approx 130$ meV; hence, one expects a smaller gap in the homogeneous limit. However, we found that the local enhancement of $J$ due to oxygen dopants restores the pairing strength and results in a pairing gap comparable to the STM data, as will be demonstrated in section 4.

In equation (2), the electron operator has been decomposed into the spin (spinon) and charge (holon) components, $c_{i\sigma} = f_{i\sigma} (b_i^\sigma)^\dagger$, $n^f_{i\sigma} = f_{i\sigma}^\dagger f_{i\sigma}$ and $n^b_i = b_i^\dagger b_i$ represent the spinon and holon densities, respectively. In the homogeneous case, the non-double occupancy constraint $\sum_{\sigma} f_{i\sigma}^\dagger f_{i\sigma} + b_i^\dagger b_i = 1$ can be satisfied by adjusting spinon and holon chemical potential independently, so the Lagrangian multiplier $\lambda_i$ is practically irrelevant. We apply the following mean field decoupling to the Heisenberg term:

$$
\tilde{S}_i \cdot \tilde{S}_j = S^z_i S^z_j - \frac{1}{2} D^\dagger_{ij} D_{ij} + \frac{1}{2} \sum_{\sigma} n^f_{i\sigma} n^f_{j\sigma}, 
$$

(3)

where

$$
\tilde{S}_i = \frac{1}{2} f_{i\alpha}^\dagger \sigma_{\alpha\beta} f_{i\beta},
$$

$$
D^\dagger_{ij} = \sum_{\sigma} f_{i\sigma}^\dagger f_{j\sigma}^\dagger.
$$

(4)
The Hamiltonian is then
\[ H = \sum_{i,j} \sigma \left( \sum_{\sigma} -t_{ij} b_i^\dagger f_{j\sigma}^\alpha f_{j\sigma} + \frac{1}{2} J S_i^z S_j^z - \frac{1}{2} D_{ij}^\alpha D_{ij} \right) - \mu_f \sum_{i,\sigma} n_f^\sigma - \mu_b \sum_i \sum_{\sigma} \lambda_i \left( n_f^\sigma + n_b^\sigma - 1 \right). \] (5)

Note that the spinon density–density term \( n_f^\sigma n_f^\sigma / 4 \) is canceled. The resulting Hamiltonian is slightly different from other slave boson decoupling schemes [20, 21]. However, we do emphasize the importance of keeping the \( S_i^z S_j^z \) term. A mean-field decoupling of this kind shows that \( S_i^z S_j^z \) accounts for the magnetic correlations that cause impurity-induced magnetization in the normal state [14]. Evidence from neutron scattering indicates an induced magnetization in the center of a vortex [22], which suggests that magnetic correlations also exist in the superconducting state. We therefore choose to adopt the mean-field decoupling in equation (3), as it is crucial for other calculations involving magnetic field.

Renormalization of the holon bandwidth is associated with the following operator:
\[ K_{ij\sigma} = f_{i\sigma}^\dagger f_{j\sigma}, \] (6)
while the renormalization of the spinon bandwidth is simply associated with \( b_i \) at low temperature. In the mean-field treatment, operators in equations (4) and (6) are replaced by their ensemble average
\[ \langle S_i^z \rangle = m_i, \]
\[ \langle D_{ij}^\alpha \rangle = d_{ij}^\alpha, \]
\[ \langle K_{ij\sigma} \rangle = \chi_{ij\sigma}, \]
\[ \langle b_i^\dagger \rangle = b_i^\dagger, \]
\[ \langle b_i^\dagger b_j \rangle = Q_{ij}. \] (7)
The nonzero value of \( \langle b_i^\dagger \rangle \) in equation (7) implies Bose–Einstein condensation (BEC) in the holon sector. At temperature relevant to the STM measurement that we primarily compare with [5], roughly \( T_{\text{STM}} < 50 \text{K} \), we find that the holon ground state takes more than 99% of the hole population at any doping larger than 5%. Thus, one can safely assume that bosonic order parameters are entirely determined by ground state wave functions, and we expect \( \langle b_i^\dagger b_j \rangle = \langle b_i^\dagger \rangle \langle b_j \rangle = Q_{ij} \) (see below). The usual mean-field procedure is applied to the above formalism [21], which yields the following Lagrangian for spinons and for bosonic excitations:
\[ L = \sum_{i,\sigma} f_{i\sigma}^\alpha \left( \partial_t - \mu_f \right) f_{i\sigma} + \sum_i b_i^\dagger \left( \partial_t - \mu_b \right) b_i + \sum_i \sum_{\sigma} \lambda_i \left( \sum_{\sigma} f_{i\sigma}^\alpha f_{i\sigma} + b_i^\dagger b_i - 1 \right) + \sum_{i,j} J S_i^z m_j \]
\[ + \sum_{i,j} -t_{ij} Q_{ij} f_{i\sigma}^\alpha f_{j\sigma} + \sum_{i,j} \left( -t_{ij} \sum_{\sigma} \chi_{ij\sigma} \right) b_i^\dagger b_j + \sum_{i,j} -\frac{J}{2} d_{ij}^\alpha D_{ij} + \text{h.c.} \] (8)
In the homogeneous d-wave superconducting state, the following ansatz is made for the order parameters:

\[
\begin{align*}
\chi_{\langle ij \rangle \sigma} &= \chi_{\langle ij \rangle \sigma}^* = \chi^0, \\
\chi_{\langle\langle ij \rangle \rangle \sigma} &= \chi_{\langle\langle ij \rangle \rangle \sigma}^* = \chi^0', \\
Q_{\langle ij \rangle \sigma} &= Q_{\langle ij \rangle \sigma}^* = Q^0, \\
Q_{\langle\langle ij \rangle \rangle \sigma} &= Q_{\langle\langle ij \rangle \rangle \sigma}^* = Q^0', \\
d_{ij} &= d_{ij}^* = \\
&\begin{cases}
  d_0 & \text{for } j = i \pm \hat{x}, \\
  -d_0 & \text{for } j = i \pm \hat{y},
\end{cases}
m_i = 0,
\end{align*}
\]

(9)

where \( \langle ij \rangle \) and \( \langle\langle ij \rangle \rangle \) denote nearest- and next-nearest-neighbor bonds, respectively. The Lagrangian of the holon and spinon are then separated. The spinon part has dispersion and d-wave gap given by

\[
\begin{align*}
\xi_k &= -2t Q^0 (\cos k_x + \cos k_y) - 4t' Q^0' \cos k_x \cos k_y - \mu_f, \\
\Delta_k &= J d_0 (\cos k_x - \cos k_y), \\
H_f &= \sum_k \Phi^*_k \begin{pmatrix} \xi_k & \Delta_k \\ \Delta_k & -\xi_k \end{pmatrix} \Phi_k, \\
\Phi_k &= \begin{pmatrix} f_{k\uparrow} \\ f_{-k\downarrow}^* \end{pmatrix}.
\end{align*}
\]

(10)

For the holon part, the Hamiltonian is

\[
H_b = \sum_k \omega_k b_k^* b_k.
\]

(11)

The order parameters are solved self-consistently by means of the Bogoliubov–de Gennes (BDG) equation with the inclusion of BEC of holons. For the spinon sector, we apply the following spin-generalized Bogoliubov transformation:

\[
\begin{align*}
&f_{i\uparrow} = \sum_n u_{n,i\uparrow} Y_n\uparrow + v_{n,i\uparrow}^* Y_n\downarrow, \\
f_{i\downarrow} = \sum_n u_{n,i\downarrow} Y_n\downarrow + v_{n,i\downarrow}^* Y_n\uparrow.
\end{align*}
\]

(12)

After diagonalization, the spinon order parameters are calculated by wave functions \( u_{n,i\sigma} \) and \( v_{n,i\sigma} \), and the Fermi distribution \( f(E_{\sigma\sigma}) \)

\[
\begin{align*}
\langle f_{i\sigma}^* f_{j\sigma} \rangle &= \sum_{n>0} \left[ u_{n,i\sigma}^* u_{n,j\sigma} f(E_{\sigma\sigma}) + v_{n,i\sigma}^* v_{n,j\sigma} (1 - f(E_{\sigma\sigma})) \right], \\
\langle f_{i\uparrow} f_{i\downarrow} \rangle &= \sum_{n>0} \left[ u_{n,i\uparrow} v_{n,i\downarrow} (1 - f(E_{\uparrow\uparrow})) + v_{n,i\uparrow}^* u_{n,i\downarrow} (1 - f(E_{\uparrow\downarrow})) \right].
\end{align*}
\]

(13)
Bosonic order parameters under BEC are calculated by assuming that all holons are condensed in the ground state

\[ \langle b_i \rangle = \sqrt{N_0 \alpha_{0,i}} , \]

(14)

where \( \alpha_{0,i} \) is the ground state wave function of equation (11), and \( N_0 = x L^2 \) is the total number of holes on an \( L \times L \) lattice. The d-wave superconducting state is characterized by the region in the phase diagram where \( \Delta_0 \neq 0 \) and \( T < T_{\text{BEC}} \). In reality, BEC in the slave boson formalism is a rather complicated issue. As pointed out in [21], if one naively adopts the BEC formalism for two-dimensional (2D) interacting bose systems, the condensation temperature is as high as \( T_{\text{BEC}} \sim O(1000) \) K near optimal doping, which is certainly unrealistic. Such a high condensation temperature is properly reproduced if one generalizes the BEC treatment, equations (7) and (14), to finite temperature. To reduce the condensation temperature to a realistic scale, one needs to include the gauge degrees of freedom and their scattering with holons [21]. Nevertheless, we are primarily interested in low-temperature STM results deep inside the condensate, \( T \ll T_{\text{BEC}} \). Therefore, the precise value of \( T_{\text{BEC}} \) does not affect the results in this paper, and our assumption that holons occupy only the ground state, equation (14), is adequate. For the calculation of gap inhomogeneity, we choose \( T = 50 \) K such that it sits reasonably below \( T_c \) in most of the doping regime, while safely above the temperature limit due to finite cluster size. STM data at this temperature are also available to compare with.

Figure 2 shows the pairing gap and DOS as a function of hole doping. Note that since holons remain condensed, STM effectively measures the spinon spectrum. The implication of figure 2 is straightforward: as doping increases, the spinon bandwidth enlarges, causing a reduction of DOS at the Fermi surface, hence a smaller pairing gap. The enlargement of the spinon band is due to a larger \( Q_0 \). This is because at low temperatures, \( Q_0 \approx n^h = x \), so the spinon bandwidth is roughly proportional to doping. Our simulation simply indicates that, although the pairing interaction is enhanced locally around each oxygen dopant, the fact that an oxygen dopant donates holes into the CuO\(_2\) plane can change the spinon DOS dramatically. This serves as another mechanism to determine the pairing gap in a larger length scale, as long as holes are practically not localized. The precise simulation that contains both local and global effects is presented in the next section.

Note that in order to compare our simulation with real cuprates, the hole doping level has to be rescaled. It is known that slave boson mean-field theory of this kind has problems in getting both temperature and doping scale correctly [23, 24]. We choose our parameters such that the simulation yields a pairing gap close to the value indicated by STM (\( \Delta \sim 20\) meV at optimal doping). The doping level is then rescaled empirically by identifying the doping where superconductivity vanishes \( x \approx 0.6 \) with that of a real cuprate \( p \approx 0.3 \); therefore

\[ x \approx 2p . \]

(15)

Throughout this paper, we denote the numerical doping scale by \( x \), and the true doping scale in cuprates by \( p \). One should rather regard equation (15) as an empirical formula, since the precise correspondence between \( x \) and \( p \) depends on the mean-field decoupling equation (3). Nevertheless, the pairing gap and critical temperature at optimal doping (\( T_c \approx 100 \) K at \( p = 0.15 = x/2 \)) is very close to the true value in BSCCO.
In this section, we incorporate oxygen dopants into the slave boson mean-field theory studied in section 3 to simulate the surface of BSCCO. We adopt the phenomenological treatment of [7] (see also [17]) that assumes that oxygen dopants locally enhance pairing strength and attract holes to their vicinity. In contrast to the weak coupling theory in section 2, enhancement of the pairing interaction is described by locally varying $J_{ij}$ in the $t-t'-J$ model,

$$J_{ij} = J + \delta J \left( n_i + n_j \right) / 2,$$

(16)
and the Coulomb interaction yields an additional term in the Hamiltonian described by locally varying potential $V_i$

$$V_i = V_0 h_i,$$

$$h_i = \sum_s \frac{\exp (-r_{is}/\lambda)}{r_{is}},$$

where $s$ is the projected position of the oxygen dopants into the plane. The precise value of $\delta J$, $V_0$, and $\lambda$ will be addressed later in the discussion of different energy and length scales. With all these effects together, the Hamiltonian in the slave boson language reads

$$H_f = \sum_{i,j} -t_{ij} Q_{ij}^+ f^\dagger_{j\sigma} f_{i\sigma} + \sum_{i,j} -t_{ij} Q_{ij}^+ f^\dagger_{j\sigma} f_{i\sigma} + \sum_{i,j} J_{ij} S^z_i m_j + \sum_{i,j} -\frac{J_{ij}}{2} d^\dagger_{ij} d_{ij} + \sum_{i,j} -\frac{\mu_f + \lambda_i}{2} n^f_{i\sigma},$$

$$H_b = \sum_{i,j} -t_{ij} \chi^*_{ij\sigma} b^\dagger_{i\sigma} b_{j\sigma} + \sum_{i,j} -t_{ij} \chi^*_{ij\sigma} b^\dagger_{i\sigma} b_{j\sigma} + \sum_{i} (V_i - \mu_b + \lambda_i) n^b_i.$$

Note that $V_i$ is present only in the bosonic sector, as it affects only charge degrees of freedom in the problem. In the disorder case, $\lambda_i$ serves as a local correction to the chemical potential in order to satisfy the nondouble occupancy constraint. To make a comparison with the density inhomogeneity in [25], we assume that oxygens are located on top of four adjacent Cu sites, although this is slightly different from the most energetically favorable position [19]. We assume that the positions of oxygen dopants are truly random, unlike the nonadjacent condition used in [25]. This is because if the oxygens are assumed to be nonadjacent to each other, then in a 2D square lattice their concentration can be at most 25%, whereas a higher oxygen concentration is necessary to achieve higher doping level in this work.

Figure 3 shows the probability distribution $P(\Delta)$ and the real space map of the pairing gap. In figure 3(a), one sees a fairly broad distribution for the doping regime $0.1 \leq x \leq 0.5$, whose mean shifts to smaller value as doping increases. This is a clear indication that the average pairing gap decreases with increasing doping, consistent with the observation in STM. On comparing figure 3(b), where the average pairing gap $\langle \Delta \rangle$ is quantified for each probability distribution, with the homogeneous gap shown in figure 2, it is evident that this $\langle \Delta \rangle$ versus $x$ behavior is inherent from the doping dependence in the homogeneous case. Since the lineshape of $P(\Delta)$ is rather peculiar, we calculate its linewidth by means of standard deviation, as shown in figure 3(c). In comparison with the STM data [5], where the linewidth narrows significantly, about 60% from the underdoped to the overdoped regimes, our simulation shows a narrowing by only about 20%. We suspect that to capture correctly this 60% reduction of $P(\Delta)$ linewidth, one needs a fine mechanism to control the overlap between superconducting islands as doping changes, for instance a fine-tuning of $\lambda$, or using a more sophisticated impurity model than equations (16) and (17). Nevertheless, our calculation demonstrates that within a simplistic impurity model with only one length scale $\lambda$ associated with impurity, we can qualitatively capture the pairing gap inhomogeneity induced by oxygens. Note that the linewidth of $P(\Delta)$ increases again in the highly overdoped case $x = 0.5$. This is because in the absence of oxygen disorder, the homogeneous gap is zero at this doping. Randomly distributed oxygens locally enhance the gap, which causes the distribution to range broadly from zero to $\Delta \approx 0.08$. 

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**Figure 3.** Simulation of BSCCO surface with interstitial oxygen dopants. The homogeneous parameters \(t, t',\) and \(J\) are the same as in figure 2, and disorder parameters are \(V_0/t = 0.2,\) \(\delta J/t = 0.18,\) \(\lambda = 0.5a.\) (a) Probability distribution \(P(\Delta)\) of the pairing gap at different doping, in a \(30 \times 30\) cluster averaged over 40 impurity configurations. (b) The average gap and (c) the standard deviation calculated from \(P(\Delta).\) (d)–(g) Real space gap map for a specific impurity configuration in different doping levels. Oxygen positions are indicated by white dots. The temperature is fixed at \(T = 50\) K in this figure.

From the real space gap map, figures 3(d)–(g), the inhomogeneity induced by oxygen dopants (white dots) is clearly visible. For each doping level, the pairing gap is locally enhanced near oxygen dopants, consistent with the weak coupling approach [7]. This is particularly evident at low doping level, as shown by red spots in figure 3(d) for \(x = 0.1,\) where a large gap region coincides with the oxygen positions. The oxygen-rich and oxygen-poor regions also give a double line structure to the gap distribution shown in figure 3(a), particularly for low doping lines from \(x = 0.1\) to 0.3. At higher doping, puddles of the large gap region start overlapping, which eventually smears out the double line structure at about \(x = 0.4.\)

We now discuss the robustness of the simulation against variation of the parameters, especially the interaction strength \(\delta J\) and \(V_0\) and the interaction range \(\lambda.\) We find that in order to have an average pairing gap decreasing with doping, \(\lambda\) has to be relatively small, roughly \(\lambda \leq 1.\) In other words, the interaction has to be modulated on an atomic scale [7]. This can be understood simply because, had the pairing enhancement due to oxygen dopants been longer range, it would have overcome the correlation-induced doping effect and enhanced the gap globally. Secondly, in order to produce a probability distribution that has comparable linewidth to STM data, the pairing enhancement must be roughly the same as the homogeneous superexchange interaction \(\delta J \approx J.\) This gives an effective coupling \(1.34J\) on the nearest \(\text{Cu–Cu}\) bonds of an oxygen dopant. The strength of Coulomb interaction \(V_0\) is also crucial for the pairing gap. We find that if \(V_0\) is too large, the holon density is highly accumulated near
oxygen dopants, causing a very low spinon density and a smaller gap, even though pairing interaction is enhanced. As also found by Nunner et al., a large value of $V_0$ can destroy the anticorrelation between the coherence peak width and position. This analysis sets up an upper limit for the Coulomb interaction $V_0 \lesssim 0.4$. We choose $V_0 = 0.2$, close to the value at optimal doping obtained by the estimation of the dielectric constant [25], although the value here is taken to be doping independent.

A simulation of the LDOS and order parameters for a specific disorder configuration is shown in figure 4. For any doping level examined, the height of the coherence peak is found to decrease with gap size and vice versa, coinciding with [7] and the STM data. Comparing the $\Delta_{ij}$ map in figure 4(f) with the $Q_{ij}$ map in figure 4(g), it is evident that regions with large gap also have large $Q_{ij}$. Similar to the homogeneous case, this is because at relatively low temperatures, bosons are fairly condensed, so $Q_{ij} \approx \sqrt{n_i^H(n_j^S)}$. Thus $Q_{ij}$ is large near oxygen dopants because the holon density is rich. Since $Q_{ij}$ enters the spinon Hamiltonian equation (18) as hopping amplitude, enhancement of $Q_{ij}$ means that the spinon hopping amplitude is locally enhanced.
around oxygen dopants, as assumed phenomenologically in section 2. On the other hand, the LDOS near the oxygen dopants has a very pronounced impurity bound state at band edge, which further reduces the coherence peak. From the phenomenological model in section 2, it is clear that this bound state is a result of hopping enhancement, i.e. larger $Q_{ij}$ near oxygen dopants. Therefore from this calculation of the LDOS, one can see the importance of strong correlations for local bandwidth and how it influences the shape of LDOS.

Figure 4(e) shows a very significant holon density inhomogeneity. This compares well with [25], where inhomogeneous charge density on the BSCCO surface was studied by means of small hole pockets. For average doping $x = \langle n^b_i \rangle = 0.33$, the holon density varies from $0.3 < n^b_i < 0.4$. However, in the weak coupling mean-field approach of [7], the hole density inhomogeneity is very moderate, roughly less than 5%. This can be understood as the following: in both [25] and the present work, charge degrees of freedom are carried by dilute holes, which screen Coulomb impurities poorly, while in [7] it is carried out by electrons, which screen Coulomb impurities very efficiently. We anticipate that an approach like that in the present work will be particularly accurate for underdoped to optimally doped cuprates, where hole density inhomogeneity is due to poor screening of dilute holes. On the overdoped side, where the system has presumably crossed over to a Fermi liquid, one should perhaps adopt weak coupling approaches like in [7] to describe accurately the screening by electrons. Note that the density inhomogeneity in the present work is slightly smaller than in [25], due to the coupling between charge and spin degrees of freedom. This is similar to a previous paper where no clear Friedel density oscillation is seen near a point-like repulsive impurity [14]. This is simply because holons, which respond to the Coulomb interaction of disorder, do not have a characteristic length scale, so its density profile is rather smooth. Due to the non-double occupancy constraint, the amplitude of charge inhomogeneity is also suppressed, because holon density is subject to the spinon density that does not respond to the Coulomb interaction directly.

Lastly, we address the issue of various correlation functions related to STM. We define the correlation functions according to [25], which is consistent with [4] but defined on a discrete lattice. The results for various correlators are shown in figure 5. The oxygen dopants are positively correlated with pairing gap. The value of correlation extrapolates to $\approx 0.7$ for

![Figure 5. Various correlation functions calculated according to [25], at doping level $x = 0.3$ and $T = 50$ K.](http://www.njp.org/)
onsite correlation $R = 0$, close to the value $\approx 0.4$ extracted from STM [4]. The holon–oxygen correlation, which is exactly the negative of spinon–oxygen correlation due to the no-double-occupancy constraint, is always positive, indicating the fact that an oxygen dopant attracts holes to its vicinity. The length scale at which correlation drops to zero is roughly $2a$, which is smaller than that reported in [25]. Note that in [7], the potential length scale $\lambda = 0.5$ yields a correlation length in good agreement with the STM results, whereas the same value of $\lambda$ gives a smaller correlation length in the current approach. We suspect that this is because spinon correlation functions are subject to holon correlation functions, and the no-double-occupancy constraint reduces the correlation length. Since $\lambda$ is treated phenomenologically, in principle it can be adjusted to yield a longer correlation length. However, in the present approach—where the oxygen concentration is rather high—our choice of a smaller $\lambda$ allows us to keep the pairing enhancement local.

5. Conclusion

We have presented a slave boson mean-field approach to compare the possible physical effects induced locally by oxygen dopants near the surface of BSCCO, and show how their interplay can explain the pairing gap revealed by STM experiments. Firstly, the enhancement of the pairing gap and the reduction of coherence peak near oxygen dopants are attributed to locally enhanced pairing interactions via superexchange. Secondly, the average spinon DOS at the chemical potential is reduced at large doping, causing the average gap to decrease, although it remains locally enhanced around oxygen dopants. We also found that since holes are attracted to the vicinity of oxygen dopants, an impurity bound state is formed at the edge of the spinon band, which causes further reduction of coherence peak, yielding a LDOS shape consistent with STM.

The present work also gives important evidence regarding various length scales and interaction strengths in the cuprates. Since the pairing gap is locally enhanced around oxygen dopants but has an average value that decreases with doping, we conclude that the length scale at which the pairing interaction is enhanced is very small $\lambda \approx 0.5a$, although the enhancement $\delta J$ is of about the same magnitude as the bare pairing interaction $J$, such that the it gives a significant gap inhomogeneity. On the other hand, the Coulomb interaction associated with the oxygen dopants is relatively small compared to the hopping amplitude, such that the accumulation of holes around oxygen dopants does not lead to a reduction of the pairing gap near oxygen dopants, although it is large enough to yield a significant charge inhomogeneity in the underdoped regime.

Acknowledgments

We thank G Khaliullin, M Mori, O P Sushkov, C Hamer, J Oitmaa and J C Davis for stimulating discussions. PJH acknowledges support from NSF-DMR-1005625. The numerical calculation in this work was performed at the facilities in the Australian National Computational Infrastructure under project no. u66.

References

[1] Howald C, Fournier P and Kapitulnik A 2001 Phys. Rev. B 64 100504
[2] Pan S H et al 2001 Nature 413 282

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[3] Lang K M et al 2002 Nature 415 412
[4] McElroy K, Lee J, Slezak J A, Lee D-H, Eisaki H, Uchida S and Davis J C 2005 Science 309 1048
[5] Gomes K K, Pasupathy A N, Pushp A, Ono S, Ando Y and Yazdani A 2007 Nature 447 569
[6] Pasupathy A N, Pushp A, Gomes K K, Parker C V, Wen J, Xu Z, Gu G, Ono S, Ando Y and Yazdani A 2008 Science 320 196
[7] Nunner T S, Andersen B M, Melikyan A and Hirschfeld P J 2005 Phys. Rev. Lett. 95 177003
[8] Maška M M, Śledź Z, Czajka K and Mierzejewski M 2007 Phys. Rev. Lett. 99 147006
[9] Foyevtsova K, Valentí R and Hirschfeld P J 2009 Phys. Rev. B 79 144424
[10] Johnston S, Vernay F and Devereaux T P 2009 Europhys. Lett. 86 37007
[11] Foyevtsova K, Kandpal H C, Jeschke H O, Graser S, Cheng H-P, Valentí R and Hirschfeld P J 2010 Phys. Rev. B 82 054514
[12] Mori M, Khaliullin G, Tohyama T and Maekawa S 2008 Phys. Rev. Lett. 101 247003
[13] Khaliullin G, Mori M, Tohyama T and Maekawa S 2010 Phys. Rev. Lett. 105 257005
[14] Gabay M, Semel E, Hirschfeld P J and Chen W 2008 Phys. Rev. B 77 165110
[15] Wang Z, Engelbrecht J R, Wang S, Ding H and Pan S H 2002 Phys. Rev. B 65 064509
[16] Zhou S, Ding H and Wang Z 2007 Phys. Rev. Lett. 98 076401
[17] Zhu J-X 2005 arXiv:cond-mat/0508646
[18] Fang A C, Capriotti L, Scalapino D J, Kivelson S A, Kaneko N, Greven M and Kapitulnik A 2006 Phys. Rev. Lett. 96 017007
[19] He Y, Nunner T S, Hirschfeld P J and Cheng H-P 2006 Phys. Rev. Lett. 96 197002
[20] Ubbens M U and Lee P A 1992 Phys. Rev. B 46 8434
[21] Lee P A and Nagaosa N 1992 Phys. Rev. B 46 5621
[22] Lake B et al 2002 Nature 415 299
[23] Ruckenstein A E, Hirschfeld P J and Appel J. 1987 Phys. Rev. B 36 857
[24] Kotliar G and Liu J 1988 Phys. Rev. B 38 5142
[25] Chen W, Khaliullin G and Sushkov O P 2011 Phys. Rev. B 83 064514