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Simulating superconducting properties of overdoped cuprates: The role of inhomogeneity

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Theoretical studies of disordered \(d\)-wave superconductors have focused, with a few exceptions, on optimally doped models with strong scatterers. Addressing recent controversies about the nature of the overdoped cuprates, however, requires studies of the weaker scattering associated with dopant atoms. Here we study simple models of such systems in the self-consistent Bogoliubov-de Gennes (BdG) framework, and compare to disorder-averaged results using the self-consistent \(T\)-matrix-approximation (SCTMA). Despite surprisingly linear in energy behavior of the low-energy density of states even for quite disordered systems, the superfluid density in such cases retains a quadratic low-temperature variation of the penetration depth, unlike other BdG results reported recently. We trace the discrepancy to smaller effective system size employed in that work. Overall, the SCTMA performs remarkably well, with the exception of highly disordered systems with strongly suppressed superfluid density. We explore this interesting region where gap inhomogeneity dominates measured superconducting properties, and compare with overdoped cuprates.

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I. INTRODUCTION

Many phenomena observed in underdoped cuprates are currently highly debated, including the origin of the so-called competing order phases: Charge and spin ordering, pseudogap, etc. Overdoped cuprates, which used to be regarded as relatively simpler, have recently attracted considerable interest, due to a few striking observations of unexpected behavior apparently inconsistent with the Landau-BCS paradigm [1–3]. One observation that has received a great deal of attention is the 2D superfluid density in overdoped \(La_{2-x}Sr_xCuO_4\) (LSCO). The Brookhaven group prepared a large sequence of high-quality, epitaxially grown overdoped films [1], and showed that in a broad range of doping, the superfluid density decreases linearly as temperature is increased. The authors also found that the zero-temperature superfluid density \(\rho_s\) at various dopings is proportional to the critical temperature \(T_c\), in contrast to BCS theory, which states that the zero temperature \(\rho_s\) should not depend on \(T_c\) at all in a clean system, but instead be simply proportional to the carrier density (which increases as the system is doped). Although this quasiproporportionality of \(\rho_s(0)\) and \(T_c\) can occur in the dirty limit [4], the authors ruled out an explanation based on disorder, since the expectation is that the linear temperature variation of the clean superfluid density is replaced by quadratic behavior in a dirty system [5], which was not observed in their films down to the lowest measurement temperature of about 3K.

In a series of papers [6–9] based on the so-called “dirty \(d\)-wave” theory, David Broun and collaborators, including one of the present authors, have argued that these data can be explained naturally and simply by accounting for the weak scattering nature of the dopant impurities, located away from the \(CuO_2\) planes, as well as the correct band structure of LSCO. In this approach, the disorder averaged self-energy of a \(d\)-wave superconductor is calculated using the self-consistent \(T\)-matrix approximation (SCTMA). These authors showed that a very good accounting of the unusual behavior of the overdoped films could be obtained within this Landau-BCS paradigm, without resorting to more exotic explanations.

The SCTMA assumes that disorder is distributed randomly, and replaces the dirty superconductor with an effective translationally invariant medium with dissipation. However, there is considerable evidence that at least some overdoped cuprate samples are quite inhomogeneous [10,11] at the nanoscale. The origin of this inhomogeneity is not completely clear. It may arise in the sample fabrication and annealing process from chemical barriers, in which case more refined annealing protocols might remove much of the inhomogeneity. Such an explanation is indeed supported by the claimed homogeneity of the epitaxially grown superconducting films [1] relative to the samples in Refs. [10,11].

On the other hand, it is possible that some of the inhomogeneity is emergent, i.e., arises in any short coherence length \(d\)-wave system with random disorder. Recently, Li et al. [3] studied the superfluid density versus doping for a model of a \(d\)-wave superconductor using BdG simulations, showed that very inhomogeneous gap distributions occurred for large concentrations of weak-to-intermediate impurity potentials, and proposed that an emergent network of \(d\)-wave superconducting islands connected by weak Josephson links was present in these samples. Such a description of the superconducting state would perforce lead to large phase fluctuations, which are accounted for neither in the SCTMA nor the BdG calculations,
but which might give rise to a low-temperature linear superfluid density behavior [12]. Interestingly, in the supplementary material of the Li et al. paper [3], the superfluid density was calculated without phase fluctuations, and nevertheless shown to be remarkably linear in $T$ even for substantial concentrations of moderate strength impurities, which is surprising, given the expectations from the dirty $d$-wave theory.

In this paper, we compare the results of the disorder averaged theory of a $d$-wave superconductor with the self-consistent BdG method. The latter is subject to finite-size effects that need to be carefully controlled, but includes both (a) scattering processes not included in the SCTMA and (b) the effects of inhomogeneity. Our goals are to test the validity of the SCTMA, which has proven so successful in explaining the properties of the overdoped cuprates [6–9], and to check the predictions of Li et al. [3] to see if alternative disorder/inhomogeneity based explanations are possible. A recent paper by Breio et al. [13] investigated some of the latter issues, and concluded on the basis of similar BdG calculations that an effective model of weakly coupled $d$-wave grains did not emerge naturally from the highly disordered regime. Here we study density of states, superconducting order parameter, and superfluid density of a disordered $d$-wave superconductor, and show that the SCTMA is remarkably robust in explaining the regime of intermediate strength disorder appropriate for dopant atoms, until regions of vanishing superconductivity begin to appear in the sample, either for very high disorder levels or very close to the critical temperature.

The quasiparticle density of states in this regime is found to be remarkably linear at low energies until the gap is nearly filled. Despite this, the penetration depth retains its quadratic behavior for disorder levels that strongly suppress the zero temperature superfluid density. This result, which contradicts that of Li et al. [3], is obtained properly only for sufficiently large system size and configuration averaging. We note their result for the $T$ dependence of $\rho_s$ was not one of the main points of their paper, nor does this remark affect their main conclusions; however it is interesting and reassuring that the expected results from the SCTMA are indeed accurate. We note further that the results reported in the present work do not contradict the result of Ref. [6], which stated clearly that the quasilinear behavior in $\rho_s(T)$ was due in large part to the particular electronic structure of LSCO, which we do not consider here.

II. MODEL

We consider a Hamiltonian for disordered cuprates which assumes weak pointlike potential impurities of a fixed strength randomly at different lattice sites. Following Ref. [3], in order to avoid issues related to the pseudogap, the homogeneous system is taken at optimal doping of $p = 15\%$, corresponding to $n = 0.85$ electron density. Disorder introduces an additional $n_{\text{imp}}$ impurities, each of which dopes the system with approximately one electron, such that $p = 0.15 + 0.5n_{\text{imp}}$. The Hamiltonian is given by

$$\hat{H} = \sum_{ij\sigma} [t_{ij} + (w_i - \mu)\delta_{ij}]c_{i\sigma}^\dagger c_{j\sigma} + \hat{H}_{\text{pair}},$$  \hspace{1cm} (1)

where $t_{ij}$ are the tight-binding hopping integrals, $w_i$ are impurity potentials at site $i$ and $\mu$ is the chemical potential. The pairing is treated in a mean-field approximation,

$$\hat{H}_{\text{pair}} = \sum_{\langle i,j \rangle} \{\Delta_{ij}c_{i\uparrow}^\dagger c_{j\downarrow} + H.c.\}$$  \hspace{1cm} (2)

with $\Delta_{ij} = V\langle c_i^\alpha c_j^\beta \rangle$ as the order parameter. A local $d$-wave order parameter $\Delta_i$ can be defined as $\Delta_i = \frac{1}{4} (\Delta_{i,i+\hat{x}} + \Delta_{i,i+\hat{y}} - \Delta_{i+\hat{x},i} - \Delta_{i+\hat{y},i})$. We take $t_{ij} = -1$ for nearest neighbor $i,j$; $t_{ij} = 0.35$ for next nearest neighbor $i,j$; and zero otherwise, corresponding to a momentum space band for the clean system of $\varepsilon_k = 2t (\cos k_x + \cos k_y) + 4t' \cos k_x \cos k_y$. This ratio of $t/t'$ is quite typical for cuprate band structures. The pairing interaction is taken as $V = 0.8t$. In the clean limit, this gives a superconducting transition temperature $kT_c = 0.085t$ when the electron density is $n = 0.85$ per unit cell. This electron density corresponds to a hole density $p = 1 - n = 0.15$ that is commonly associated with optimal doping in cuprates. The corresponding $d$-wave gap function is $\Delta_k = \Delta_0 (\cos k_x - \cos k_y)$, with $\Delta_0 = 2\Delta_i = 0.0935t$. Note that since in cuprates $t \sim 300$ meV from ARPES, $T_c \sim 300$ K, which is artificially high. The parameters are chosen in this way to allow a direct comparison with Ref. [3], and for numerical convenience. This amounts to expressing quantities with dimensions of energy in units of $t$ without explicitly mentioning the unit.

The impurity potential $w_i$ at any site $i$ is chosen randomly to be either 0 or 1. A scattering potential of 1 ($t$) amounts to a moderate strength of the potential scatterers we are modeling, well away from the unitarity limit, but still considerably larger than the realistic ab initio dopant potentials of order 0.2$-0.3$t calculated in Ref. [9] for Sr dopants in LSCO. In this sense our study is qualitative, and makes no attempt to compare to real cuprate systems. We have also not attempted to include O vacancies and other sources of disorder discussed in Ref. [9]. The chemical potential $\mu$ is determined self-consistently via iterative diagonalization of the BdG Hamiltonian. Fixing the average electron density in the system leads to a unique chemical potential via the self-consistent diagonalization.

III. RESULTS

We work in a $40 \times 40$ lattice with periodic boundary condition, and average over several random impurity configurations at various values of average impurity density $n_{\text{imp}}$. To increase the resolution of the spectrum and for the purpose of studying the system at sufficiently low temperature, the supercell method is employed to ensure a dense set of energy levels (for details see Appendix A). With a bandwidth of $\sim 8t$ and only $40 \times 40$ levels, the level-spacing may not be sufficient at low energies if supercells are not used. Averaging over random configurations of impurities was also used to improve the statistics. Averaging over more and more impurity configurations makes the system effectively more and more translationally invariant. Typically, averaging over 40 random configurations of impurities appears to be sufficient to get an accurate low-energy behavior.

The BdG equations are iteratively and self-consistently solved for the order parameter and chemical potential. If
FIG. 1. Maps of typical random impurity configurations and corresponding superconducting d-wave order parameters $\Delta_i$ (colorscale) at three impurity concentrations: (a) 4%, (b) 12.5%, and (c) 22% at $kT/t = 0.001$. Impurity locations are indicated by red crosses. The impurity potential is $1t$.

The impurity concentration in the system is varied with fixed chemical potential, doping varies proportional to the impurity concentration, and may be compared [3] roughly to a cuprate scale as $p = 0.15 + n_{\text{imp}}/2$. Note, however, that the actual effect of doping on the electronic structure is neglected here, and the rate of doping change with impurity concentration depends in the model entirely on the potential of each individual dopant, taken here to be $w = 1t$. Results for $d$-wave order parameter, density of states and superfluid density are summarized below.

A. d-wave order parameter

First, the variation of the superconducting order parameter was studied as a function of the impurity concentration. An impurity concentration of $0 < n_{\text{imp}} < 1$ empirically corresponds to a hole doping of roughly $\Delta p \approx 0.5n_{\text{imp}}$ with respect to the optimal filling of $p = 0.15$. We worked within an impurity concentration range of $n_{\text{imp}} = 0.02-0.3$. Typical maps of impurity configurations along with the $d$-wave order parameter at moderate to high impurity concentrations are shown in Fig. 1. It is clear for the highest concentration that impurities of this strength suppress the superconductivity nearly completely, and that the resistive transition temperature, which depends on the existence of a percolating path of nonzero order, might be zero.

Note that it is known that electrostatic potentials of this kind cannot reproduce the detailed statistics of optimally doped cuprates, which require in addition a disorder component of the Andreev type [14]. Nevertheless it is clear that dopant disorder of this form can lead to extensive gap inhomogeneity for higher concentrations [3].

In these simulations, since the pairing interaction is uniform on all nearest-neighbor bonds, a small local gap component of $s$-wave symmetry will be induced by the disorder. In addition, higher concentrations will lead to time-reversal-symmetry breaking [3,13] by local currents, despite the fact that the Hamiltonian contains no terms that explicitly break time reversal symmetry. We study neither of these effects here.

The order parameter maps shown in Fig. 1 hint at another interesting aspect of the highly disordered state, namely that the order parameter inhomogeneity increases with $n_{\text{imp}}$. Figure 2(a) compares the temperature dependence of the average bond order parameter for a number of disorder configurations; each data point corresponds to the average of $\Delta_i$ over the lattice for a single configuration. Here it is seen that not
only does the gap distribution become more inhomogeneous with increasing disorder, it also becomes more sensitive to unusual local disorder configurations [3], giving rise to a much larger dispersion of average gap values over configurations. Also shown are the results for the same concentrations and model within the SCTMA (described in Appendix D). For the less disordered systems, the SCTMA describes the T-dependent order parameter semiquantitatively at low T, but predicts gap closings at lower $T_c$s than BdG (where $T_c > 0$ only means that $\Delta_1 > 0$ on as little as 1 site). For the most disordered case shown, the SCTMA predicts $\Delta = 0$ at all temperatures. The strongly inhomogeneous states found in BdG indeed persist up to much higher doping, as seen in Fig. 2(b).

**B. Density of states**

In the BCS theory of a clean $d$-wave superconductor, the low-temperature linear-$T$ term arises from the linear in $\omega$ density of states. It is therefore interesting to study how the density of states varies with energy in the disordered case with intermediate strength scattering. The single-spin local density of states (LDOS) at lattice site $i$ can be calculated from the eigenvectors $U_{is,m\pi}$ and eigenvalues $E_{ms}$ of the BdG Hamiltonian, Eq. (1),

$$\rho(r_i, \omega) = -\frac{1}{\pi} \text{Im} \left( \sum_{m,s} \frac{|U_{is,m\pi}|^2}{\omega - E_{ms} + i\eta^+} \right),$$  

where $\eta^+$ is an artificial broadening, the pair $(i, \uparrow)$ indicates the projection of the eigenstate $(n, s)$ onto lattice site $i$ with spin up. Here, the pair $(n, s)$ labels eigenstates with pseudospin $s = \uparrow, \downarrow$ such that $E_{ns} > 0$ and $E_{n\downarrow} < 0$ (see Appendix C). While there are several BdG studies with strong impurities [15–19], there are relatively few works to address weaker scatterers in the overdoped region [3,20]. From the SCTMA, it is expected that the residual value of the normal state DOS as either the concentration or the strength of impurities is increased [7]. The residual value is clearly seen in Fig. 3 to increase rapidly from zero as disorder is added.

The more interesting feature of the LDOS for four disorder concentrations shown in Fig. 3 is the persistent V-shape spectrum at low energies for the BdG calculation even with relatively high impurity concentration, where the SCTMA shows quadratic trend at low energies. In the strong-impurity case, the low-energy bound states hybridize to form a broad plateau-like impurity band, also seen qualitatively in the BdG results with the exception of an energy range exponentially small in the Dirac cone anisotropy, $\sim \exp(-E_F/\Delta_0)$ [21]. The positive energy coherence peak in the clean system is at $\omega/t \sim 0.18$ and this does not much affect at the impurity concentration of 12.5%. The peak shifts towards zero as $n_{imp}$ is further increased, however. While the peak feature at negative energies resembles a coherence peak, it is in fact the van Hove singularity in this system, and the coherence peak itself is invisible in the figure.

**C. Superfluid density**

In a tetragonal superconductor, the penetration depth for in-plane screening currents is independent of the direction of current flow within the plane. It is thus sufficient to calculate the penetration depth for the case in which the magnetic vector potential and current are both aligned with the $x$ axis, i.e., $\lambda_{xx}$. From the Kubo formula, the inverse squared penetration depth is the sum of diamagnetic and paramagnetic contributions,

$$\lambda^{-2} \equiv \lambda_{xx}^{-2} = \frac{4\pi e^2}{c^2} (K_{xx}^{dia} + K_{xx}^{para}).$$  

where $e$ and $c$ are the electron charge and speed of light, respectively. The two contributions take on well-known forms in the clean limit (Appendix B), but are less familiar in the absence of translational symmetry [22,23]. The diamagnetic response kernel is

$$K_{xx}^{dia} = \frac{1}{N} \sum_{m,s} (\tilde{M}^{-1})_{ms,ms} f(E_{ms}),$$  

where $\tilde{M}^{-1}$ is the matrix representation of the inverse effective mass in the basis of eigenstates of the BdG Hamiltonian (Appendix C) and $f(E_{ms})$ is the Fermi-Dirac function for the BdG energy eigenvalues. The paramagnetic response kernel is

$$K_{xx}^{para} = \frac{1}{N} \sum_{m,s} \sum_{x,x'} |\tilde{\zeta}_{ms,x\pi}|^2 \frac{f(E_{ms}) - f(E_{ns})}{E_{ms} - E_{ns}}.$$  

Here, $\tilde{\zeta}$ is the representation of the quasiparticle velocity in the basis of eigenstates of the BdG Hamiltonian and, as above,
m, n and \( s, s' \) are eigenenergy and pseudospin indices, respectively. \( N \) is the number of lattice points. Detailed expressions for \( \bar{M}^{-1} \) and \( \xi \) are given in Appendix C.

It is also usual to define a related quantity, the superfluid density, as

\[
\rho_s = \frac{m c^2}{4\pi e^2} \lambda^{-2}.
\]

For a parabolic band, \( m \) is the electron effective mass; otherwise, one may define \( m \) from the clean-limit expression

\[
\rho_s(n) = \frac{N_c}{2\pi^2 K_{\text{dis}}(T=0)}
\]

This choice correctly captures the fact that \( \rho_s = n \) at \( T = 0 \) in the clean limit, and is independent of either \( T_c \) or the order parameter. For the qualitative behavior (and direct comparison with results in the supplementary information of Ref. [3]), the fundamental constants have been suppressed in the plots for superfluid density.

Let us at this point connect the superfluid density at low temperatures with the density of states at low energies as discussed in Sec. III B. In BCS theory, there is a simple expression relating the superfluid density and density of states. Writing \( \rho_s = 1 - \rho_n \), with \( \rho_n \) the normal fluid density, one obtains [24]

\[
\rho_n = \int d\omega \rho(\omega) \left[ -\frac{df}{d\omega} \right].
\]

For a clean d-wave superconductor, the density of states is \( \rho(\omega) \propto \omega \), and it follows immediately that \( \rho_n \propto T \). If the density of states at low energy has a quadratic behavior, \( \rho(\omega) = \gamma + \omega^2/\Delta_0^2 \), and one simply substitutes this into the integral above, one expects that \( \rho_n(T) = \text{const.} + \frac{\pi}{2}(\pi T/\Delta_0)^2 \), such that \( \rho_s \) decreases quadratically with \( T \).

1. Zero temperature

The zero-temperature superfluid density is a fundamental measure of the strength of superconductivity. In Fig. 4, we show calculations for \( \rho_s \) from both our BdG numerical simulations and the SCTMA. As expected, the superfluid density is seen to decrease monotonically as the impurity concentration is increased in both approximations. At lower impurity concentrations, the BdG and SCTMA predictions for the suppression of \( \rho_s \) are virtually identical. It is only when the superfluid density falls to about 10% of its clean value that the two approaches differ significantly. As is well-known from previous work, \( \rho_s \) vanishes linearly with disorder within the SCTMA, as in the classic Abrikosov-Gor’kov treatment of magnetic impurities in an s-wave superconductor [25]. In the BdG case, however, \( \rho_s(0) \) has a concave-up tail that persists to high impurity concentrations, similar to what was observed in Ref. [3]. This behavior is expected, because the SCTMA describes the effects of an average order parameter, whereas in the BdG case, islands of nonzero \( \rho_s \) in rare regions favorable for superconductivity [26] persist well past the SCTMA critical concentration, and indeed well beyond the critical disorder where regions of nonzero order parameter percolate across the numerical sample (see Fig. 1).

It is also important to confirm the “non-BCS” quasiproportionality of \( \rho_s(0) \) and \( T_c \) reported in Ref. [1] and reproduced in the subsequent SCTMA study [6]. In a clean BCS superconductor, the low-temperature superfluid density is equal to the density of conduction electrons in a normal metal. When impurity scattering suppresses both \( T_c \) and \( \rho_s \) monotonically, a quasiproportionality between the two as observed in Ref. [1] can result, however. Accurate determination of \( T_c \) is not trivial in a finite size system as considered in BdG. Here to get each data point for \( T_c \) in the figure, we take the superfluid density at three highest temperature points of the corresponding \( \rho_s \) versus \( T \) data and fit a quadratic curve through those three points to estimate the \( T_c \). The result for \( T_c \) versus \( \rho_s(0) \) determined in this way is shown in Fig. 5, where we compare results for both approximations. The functional form is clearly not consistent with the BCS result \( \rho_s = n \), but neither is it consistent with the linear relation found in the experiment of Ref. [1]. We have verified that decreasing the impurity potential to correspond to the Born limit used in Ref. [6] does not affect the SCTMA curve significantly. The deviation from the quasilinear behavior is therefore due to band structure effects, specifically the proximity of the van Hove singularity to the Fermi level in the LSCO ARPES-derived band structure used in that work.

2. Temperature dependence of superfluid density

Next, we focus on the temperature dependence of the superfluid density in detail at a few specific impurity concentrations to see if the physics included in the BdG approach allows for an understanding of the approximate linear-\( T \) behavior of the penetration depth reported in Ref. [1]. We were also motivated by the BdG calculations in Ref. [3], which seemed to indicate that asymptotic linear-\( T \) behavior was often found for quite disordered samples.
FIG. 5. $T_c$ vs superfluid density $\rho_s$ at zero $T$ for BdG (blue) and SCTMA (red).

For this range of impurities, averaging was done over the $40 \times 40$ system for forty random configurations at each value of impurity concentration and $15 \times 15$ supercells were used. BdG results for the superfluid density temperature dependence, together with the equivalent result for the SCTMA, are presented in Figs. 6(a) and 6(b). The superfluid density in the clean, optimally doped cuprates typically decreases almost linearly as temperature increases [27], reflecting the linear low-energy density of states of a clean $d$-wave superconductor. However, strong disorder changes this linear-$T$ behavior into a quadratic behavior over an impurity bandwidth rather rapidly [28], as predicted by SCTMA [5,29]. Both approximations give very similar quantitative results over nearly the entire temperature range for the smaller disorder concentrations shown in Fig. 6(a), and the temperature dependencies are qualitatively similar even for larger disorder concentrations, until the disorder concentration approaches the critical disorder for the destruction of superconductivity. These discrepancies arise when the BdG superfluid density is suppressed to roughly 10% of its clean value. In overdoped LSCO, this would correspond to high levels of overdoping, near the end of the superconducting dome, e.g., for $T_c$ at or below $T_c = 10$ K [1].

In the weak scattering limit with generic bands, a larger concentration of impurities is necessary to create a clear quadratic low-$T$ behavior. From the SCTMA viewpoint, in contrast to the strong impurity case, there is no analogous impurity bandwidth over which the $T^2$ is expected. Indeed, Lee-Hone et al. [6] found nearly linear-$T$ behavior over nearly the entire doping disorder range, but emphasized that a significant role was played by the particular Fermi surface of overdoped LSCO in the intermediate-to-low temperature regime.

3. Supercell dependence of superfluid density

As mentioned above, some of the higher disorder concentrations of the $\rho_s$ versus $T$ results shown in the supplementary information of Ref. [3] appear to show low-$T$ linear behavior, in contrast to the results we have just presented. Identical band structures and disorder potentials were used in both works. To investigate possible origin of this mismatch, we studied the temperature variation of superfluid density with different number of supercells used at each of impurity concentrations 12.5%, 18.7%, and 22%. The details of the implementation are given in Appendix A. For 18.7% and 22%, we compared

FIG. 6. Effect of disorder on the temperature-dependent superfluid density. Superfluid densities at various temperatures were averaged over 40 random impurity configurations (with $15 \times 15$ supercells) at impurity concentrations (a) 0%, 4%, and 12.5%; (b) 12.5%, 18.7%, and 22%. Note that, for the last two cases, SCTMA predicts that superconductivity is entirely suppressed.
FIG. 7. Supercell dependence of superfluid density. Superfluid density averaged over 40 impurity configurations as a function of the temperature at fixed impurity concentration 12.5% with different number of supercells used. The superfluid density scale in this figure is not normalized with $\rho_s(T \to 0)$ or $\rho_{\text{Clean}}(T \to 0)$.

just between the effects of having no supercell and 15 × 15 supercells. For 12.5%, we additionally used a supercell lattice of intermediate size 5 × 5. The result is presented in Fig. 7.

It is to be emphasized that for small systems of finite size, the use of supercell lattice is quite important at low temperature to create a sufficient density of states available within a narrow energy range. Remarkably, the low-temperature behavior of superfluid density without any supercell resembles the results from Ref. [3] quite well, so we speculate that this effect is responsible for the discrepancy. The deviation from linear behavior becomes increasingly apparent in case of 18.7% and 22% impurities.

4. Superfluid density with self-consistent versus homogeneous average gap

There are several major differences between the BdG and SCTMA methods discussed here. The BdG suffers from finite size corrections that need to be carefully estimated, as shown above; however it includes both the effects of self-consistent variation of the order parameter in response to a given disorder configuration, as well as quantum interference corrections (crossing diagrams in the self-energy and vertex corrections) neglected in the one-impurity $T$ matrix. The former source of difference is easy to check by performing non-self-consistent BdG calculations with artificial homogeneous order parameter. For this, the inhomogeneous superconducting gap for each configuration was replaced with its average value at the corresponding temperature, over the lattice before calculating the superfluid density. The result is presented in Fig. 8. The self-consistent (inhomogeneous) result is seen to be smaller than the non-self-consistent (homogeneous average gap) result in the weakly disordered limit, where the superconductor is largely clean and the order parameter is suppressed only around individual impurities. On the other hand, at large disorder, the system breaks up into distinct islands of relatively large superfluid density, such that the inhomogeneous result is slightly larger. However, the differences between the homogeneous and inhomogeneous BdG results are always small. This suggests that the effects of inhomogeneity are not important to understanding the differences between the two mean-field approximations presented here, BdG and SCTMA; rather, quantum interference diagrams beyond $T$ matrix neglected in the SCTMA drive the differences at larger disorder. Homogeneity can of course become much more important to the actual superfluid density once phase fluctuations are included in the cases of very small superfluid density, however. These effects are beyond the scope of this work.

IV. DISCUSSION

In summary, we studied various superconducting properties in optimally doped to overdoped cuprates using the impurity averaged self-consistent real-space BdG formalism with supercells and periodic boundary condition. The doping and impurity parameters were taken to correspond exactly to the previous study [3] that unexpectedly found a linear $T$ dependence of the superfluid density for some highly disordered cases. The strength of the impurity assumed in that work is probably significantly higher than realistic values for real spacer layer dopants [9] and may be classified as moderate since the potential is a significant fraction of the bandwidth. Nevertheless we compared directly and found quadratic $T$ behavior for the analogous cases studied by Li et al. We have suggested here that the discrepancy lies in the use of too few states in the low-energy region of the simulations of Ref. [3].
We emphasize again, however, that these results were not the main point of their work, and our results confirm theirs in most other points of comparison.

We have shown that in this intermediate strength scattering regime, the BdG density of states shows quite remarkably robust linear low-energy excitation spectrum. Nevertheless, the low-temperature superfluid density behavior apparently displays the usual quadratic low-\(T\) temperature dependence expected for a dirty \(d\)-wave superconductor. This is an unexpected result based on the usual BCS argument, whereby the leading \(\omega\) term in \(\rho(\omega)\) near the Fermi level determines the low temperature \(T\) dependence of the superfluid density. Possible sources of this discrepancy include Andreev scattering from gap inhomogeneities, and quantum interference from multi-impurity scattering (crossing diagrams). Figure 8 shows that gap inhomogeneities do not significantly change the low-\(T\) superfluid density. We therefore conclude that the multi-impurity scattering effects are responsible for the deviation from the BCS relation between the density of states \(\rho(\omega)\) and the superfluid density \(\rho_s(T)\). Equation (9) is therefore not reliable in strongly disordered cases.

Gap inhomogeneity causes shifts in \(\rho_s\) at low \(T\). The direction of the shifts is seen in Fig. 8 to depend on the impurity concentration. For low impurity concentration, the gap inhomogeneities are isolated, and therefore lead to a small local suppression of \(\rho_s\) near each impurity. This adds an Andreev scattering channel that further suppresses the superfluid density. On the other hand, for large impurity concentrations, the inhomogeneity increases \(\rho_s\) slightly since now large islands of superconductor preferentially form in cleaner regions where ordinary impurity scattering does not suppress \(\rho_s\).

These studies suggest that the self-consistent \(T\)-matrix approximation works very well at low temperature compared to full quantum calculations for weak to intermediate strength scattering potentials, even up to quite large disorder concentrations, in contrast to strong scattering potentials [15]. Despite the fact that we found here results for the superfluid density displaying substantial quadratic behavior at low \(T\), as expected for SCTMA at high concentrations even for weak scatterers, it does not invalidate the result of Ref. [6]. This is because unlike this reference, our studies neglect the effect of the doping on the van Hove singularity near the Fermi level, as occurs in LSCO.

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APPENDIX A: SUPERCELL METHOD

The numerical calculations in this article are based on the Bogoliubov-de Gennes (BdG) method. For a homogeneous system, results from the real space BdG calculations can also be readily obtained from the corresponding momentum space equations because of translation invariance. One can freely choose as dense a momentum space grid as one wants which in real space would correspond to a bigger and bigger part of an infinite homogeneous lattice. This larger portion of lattice, however, can also be obtained from stitching the ‘lattice-portion corresponding to the coarse momentum grid’ together one after another in one or more of the spatial directions. The ‘entire lattice-portion corresponding to the coarse momentum grid’ can be called a supercell in real space and the momentum corresponding to the periodicity of these supercell is called supercell momentum. Obviously, the neighboring supercells couple with each other through the sites near their margin/boundary only, the supercell being just an imaginary construct of grouping lattice sites together. Hence, for a square lattice (which has been used in this paper), only nearest-neighbor (NN) and next-nearest-neighbor (NNN) supercells can couple with each other unless the supercell size is not critically small compared to the hopping range of the original lattice sites.

The corresponding scenario of stitching together the inhomogeneous lattices of a finite size is particularly useful for increasing smoothness of the spectrum. In this case, within the supercell there is no notion of momentum because of lack of translational invariance and the eigenstates of the inhomogeneous finite lattice are to be obtained by numerical diagonalization, often starting from the real space basis. However, subsequently, the use of many supercells effectively creates a narrow band of energy around each of these numerically obtained eigenenergies, thereby increasing the spectrum resolution.

If \(M \times M\) supercells are used, the supercell BdG equation in matrix form can be written as

\[
\begin{pmatrix}
    h(K) & \Delta(K) \\
    \Delta^T(-K) & -h^T(-K)
\end{pmatrix}
U_{nm}(K) = E_{nm}(K)U_{nm}(K)
\]

(A1)

with

\[
h(K) = \sum_{l} T_{0l} e^{K R_l},
\]

(A2)

\[
\Delta(K) = \sum_{l'} \Delta_{0l'} e^{K R_{l'}}.
\]

(A3)

Here \(K\) is supercell momentum, \(R_l\) is the coordinate of the \(l\)th supercell, \(T_{0l}\) is the supercell hopping matrix (of size \(N^2 N_o \times N^2 N_o\) where the size of the supercell is \(N \times N\) and each site has \(N_o\) orbitals) between the supercell at origin and the \(l\)th supercell, and \(\Delta_{0l}\) is the supercell gap matrix (of size \(N^2 N_o \times N^2 N_o\)) between the supercell at origin and the \(l\)th supercell. The matrix \(U\) diagonalizes the Hamiltonian and \(U_{nm}(K)\) is the \(ns\) column-vector of \(U(K)\). In this work we have \(N_o = 1\) and number of lattice points \(N = N^2\). Most of the matrix elements of \(T_{0l}\) and \(\Delta_{0l}\) (for \(l \neq 0\)) are zero unless they involve sites near the supercell-boundary which maintain the supercell-supercell coupling.

\(T_{0l}\) and \(\Delta_{0l}\) are exactly same as the hopping matrix and the gap matrix of the \(N \times N\) inhomogeneous lattice, whose spectrum is obtained by explicit iterative self-consistent diagonalization to get eigenenergies indexed by \(n\). \(K\) can take values as \(K_x, K_y = \ldots, \ldots, K_x = \ldots, K_y = \ldots\)
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where $\xi_k$ is the single particle energy with respect to chemical potential as specified in the manuscript, $E_k = \sqrt{\xi_k^2 + \Delta_k^2}$. The first sum gives the diamagnetic contribution and the second sum is the paramagnetic contribution. We have tested that the calculated penetration depth using this formulas is quite strongly dependent on the number of momentum space points. The diamagnetic term has contributions from the full Brillouin zone and the paramagnetic term has dominant contributions from the nodal areas. The diamagnetic term can be written in a different form using integration by parts, which leads to an alternate expression for the penetration depth [30]

$$\lambda_{xx}^{-2} = \frac{4\pi e^2}{c^2} \sum_k \left[ \left( \frac{\partial \xi_k}{\partial k_x} \right)^2 \left( \frac{\Delta_k}{E_k} \right)^2 - \left( \frac{\partial \xi_k}{\partial k_x} \right) \left( \frac{\partial \Delta_k}{\partial k_x} \right) \frac{\Delta_k \xi_k}{E_k^2} \right] \left[ \frac{1}{E_k} - \frac{\partial}{\partial E_k} \right] \tanh \frac{\beta E_k}{2}. \quad (B2)$$

Although mathematically equivalent to Eq. (B1), this alternate expression converges differently as a function of the number of $k$ points in the calculation. For finite systems, this can lead to inaccurate results for the superfluid density particularly at low temperatures. In the real space calculation, we are using the (effective) multiband analog of Eq. (B1) which shows a dependence on the number of supercell $K$ points especially at low temperatures, as demonstrated in Fig. 7.

**APPENDIX C: INVERSE MASS AND VELOCITY**

The Hamiltonian for an $N$ site system in the usual fermionic basis is

$$\hat{H} = c^\dagger H c \quad \text{with} \quad H = \begin{pmatrix} \frac{\hbar}{\Delta} & \Delta^* \\ \Delta & -\hbar \end{pmatrix}. \quad (C1)$$

where $H$ is a $2N \times 2N$ matrix, $\hbar$ and $\Delta$ are $N \times N$ matrices with matrix elements

$$h_{ij} = t_{ij} + (w_i - \mu) \delta_{ij}, \quad (C2)$$

$$\Delta_{ij} = V \langle c_{i\uparrow} c_{j\downarrow} \rangle, \quad (C3)$$

and $c$ and $c^\dagger$ respectively column and row vectors defined as

$$c = (c_{\uparrow1} \ldots c_{\uparrow N} c_{\downarrow1} \ldots c_{\downarrow N}). \quad (C4)$$

$H$ is diagonalized by a unitary matrix $U$ such that $U^\dagger H U = E$, with $E$ a diagonal matrix of eigenenergies. The quasiparticle operators for the diagonalized Hamiltonian are

$$\Gamma^\dagger = c^\dagger U = (\gamma_{1\uparrow} \ldots \gamma_{N\uparrow} \gamma_{1\downarrow} \ldots \gamma_{N\downarrow}) \quad (C5)$$

where in $\gamma_{ns}$, $n$ labels an eigenstate and $s$ a pseudospin. The eigenenergies satisfy $E_{ns} > 0$ and $E_{ns} < 0$.

The paramagnetic part of the electromagnetic response kernel is given by [22]

$$K_{xx}^{\text{para}} = \frac{1}{N} \sum_{m,n} \sum_{s,s'} |\tilde{\xi}_{ms,ns}|^2 \frac{f(E_{ms}) - f(E_{ns})}{E_{ms} - E_{ns}}, \quad (C6)$$

**APPENDIX B: CLEAN PENETRATION DEPTH IN REAL AND K SPACE**

It is useful for comparison to real-space BdG and SCTMA calculations to have full results for the superfluid density of the corresponding clean system, which can be calculated more accurately in momentum space for bands $\xi_k$ and gap function $\Delta_k$ as given in the main text. The result is (see, e.g., Refs. [23,30])

$$\lambda_{xx}^{-2} = \frac{4\pi e^2}{c^2} \sum_k \left( \frac{\partial \xi_k}{\partial k_x} \right)^2 \left( 1 - \frac{\xi_k}{E_k} \tanh \frac{\beta E_k}{2} \right), \quad (B1)$$

FIG. 9. Illustration of the supercell method: A supercell lattice of size $3 \times 3$ (periodically continued both horizontally and vertically) consisting of supercells of size $N \times N$ where each lattice site hosts 1 orbital. The double-way arrows show the supercell-supercell coupling in which the sites near the supercell boundaries participate.

The Hamiltonian for an $N$ site system in the usual fermionic basis is

$$\hat{H} = c^\dagger H c \quad \text{with} \quad H = \begin{pmatrix} \frac{\hbar}{\Delta} & \Delta^* \\ \Delta & -\hbar \end{pmatrix}. \quad (C1)$$

where $H$ is a $2N \times 2N$ matrix, $\hbar$ and $\Delta$ are $N \times N$ matrices with matrix elements

$$h_{ij} = t_{ij} + (w_i - \mu) \delta_{ij}, \quad (C2)$$

$$\Delta_{ij} = V \langle c_{i\uparrow} c_{j\downarrow} \rangle, \quad (C3)$$

and $c$ and $c^\dagger$ respectively column and row vectors defined as

$$c = (c_{\uparrow1} \ldots c_{\uparrow N} c_{\downarrow1} \ldots c_{\downarrow N}). \quad (C4)$$

$H$ is diagonalized by a unitary matrix $U$ such that $U^\dagger H U = E$, with $E$ a diagonal matrix of eigenenergies. The quasiparticle operators for the diagonalized Hamiltonian are

$$\Gamma^\dagger = c^\dagger U = (\gamma_{1\uparrow} \ldots \gamma_{N\uparrow} \gamma_{1\downarrow} \ldots \gamma_{N\downarrow}) \quad (C5)$$

where in $\gamma_{ns}$, $n$ labels an eigenstate and $s$ a pseudospin. The eigenenergies satisfy $E_{ns} > 0$ and $E_{ns} < 0$.

The paramagnetic part of the electromagnetic response kernel is given by [22]

$$K_{xx}^{\text{para}} = \frac{1}{N} \sum_{m,n} \sum_{s,s'} |\tilde{\xi}_{ms,ns}|^2 \frac{f(E_{ms}) - f(E_{ns})}{E_{ms} - E_{ns}}, \quad (C6)$$
where \( m, n \) are eigenenergy indices, \( s, s' \) are pseudospin indices, and

\[
\tilde{\zeta} = U^\dagger \begin{pmatrix} \zeta & 0 \\ 0 & \zeta \end{pmatrix} U
\]  

(C7)

is the representation of the electron velocity operator \( \zeta \) in the basis of eigenstates of the Hamiltonian. In the basis of lattice sites, \( \zeta \) is an \( N \times N \) matrix with matrix elements \( \zeta_{lm} = i(x_l - x_m) t_{lm} \), where \( t_{lm} \) is the tightbinding hopping matrix element between sites \( l \) and \( m \) and \( x_l \) and \( x_m \) are the \( x \) coordinates of the lattice site location. This form of the velocity operator originates in linear response theory from the Peierls substitution in the presence of a perturbing field.

The diamagnetic part of the electromagnetic response kernel is

\[
K_{xx}^\text{dia} = \frac{1}{N} \sum_{m,s} (\tilde{M}^{-1})_{ms,ms} f(E_{ms}),
\]

with

\[
\tilde{M}^{-1} = U^\dagger \begin{pmatrix} M^{-1} & 0 \\ 0 & -M^{-1} \end{pmatrix} U
\]

(C9)

and

\[
(M^{-1})_{ij} = -i t_{ij}(x_i - x_j)^2.
\]

(C10)

The inverse squared penetration depth and superfluid density are then given by

\[
\lambda_{xx}^{-2} = \frac{4 \pi e^2}{c^2} (K_{xx}^\text{dia} + K_{xx}^\text{para})
\]

(C11)

and

\[
\rho_s = \frac{mc^2}{4 \pi e^2} \lambda_{xx}^{-2}.
\]

(C12)

The formalism described in this section can be extended to the supercell method in order to facilitate numerical calculations. To this end, one needs to Fourier transform the Hamiltonian Eq. (C1) to a form like that in Eq. (A1); the velocity and mass operators can be Fourier transformed using the prescription in Eq. (A2) and acquire phase factors for displacements \( x_i - x_j \) connecting different supercells. The sums over bands in Eqs. (C6) and (C8) then pick up an additional sum over supercell momenta \( \mathbf{K} \).

**APPENDIX D: DIRTY D-WAVE THEORY**

In the presence of impurities, the superconducting Green’s function is

\[
G_{\mathbf{k}}(i\omega_n) = [i\omega_n \tau_0 - \tilde{\xi}_\mathbf{k} \tau_3 - \Delta_\mathbf{k} \tau_1 - \Sigma(i\omega_n)]^{-1},
\]

(D1)

where \( \Sigma(i\omega_n) \) is a \( 2 \times 2 \) matrix that incorporates the effects of impurity scattering. Within SCTMA, the self-energy for pointlike impurities is obtained from the \( T \) matrix via

\[
\Sigma(i\omega_n) = n_{\text{imp}} T(i\omega_n),
\]

with the \( 2 \times 2 \) \( T \)-matrix is

\[
T = [\tau_0 V^{-1} - \tau_3 g(i\omega_n)]^{-1} \tau_3
\]

(D3)

and

\[
g(i\omega_n) = \frac{1}{N} \sum_k G_{\mathbf{k}}(i\omega_n).
\]

(D4)

The impurity potential is set to \( V = 1 \). For a \( d \)-wave gap with bond order parameters \( \Delta_{\pm x} \), we have \( \Delta_\mathbf{k} = \Delta_0 \eta_\mathbf{k} \) and

\[
\Delta_0 = -\frac{J}{\beta N} \sum_{\mathbf{k}} \eta_\mathbf{k} [c_{\mathbf{k} \downarrow} c_{\mathbf{k} \uparrow}] = -\frac{J}{\beta N} \sum_{\mathbf{n},\mathbf{k}} \eta_\mathbf{k} [G_{\mathbf{k}}(i\omega_n)]_{12},
\]

with \( \eta_\mathbf{k} = \cos k_x - \cos k_y \). It is common to write

\[
\tilde{\omega}_n = \omega_n - \frac{1}{2} \text{Im} \mathcal{T} [\Sigma(i\omega_n) \tau_0],
\]

\[
\tilde{\xi}_\mathbf{k}(i\omega_n) = \tilde{\xi}_\mathbf{k} + \frac{1}{2} \text{Re} \mathcal{T} [\Sigma(i\omega_n) \tau_3],
\]

\[
\tilde{\Delta}_\mathbf{k}(i\omega_n) = \Delta_\mathbf{k} + \frac{1}{2} \text{Re} \mathcal{T} [\Sigma(i\omega_n) \tau_1],
\]

so that

\[
\Delta_0 = -\frac{J}{\beta N} \sum_{\mathbf{n},\mathbf{k}} \eta_\mathbf{k} [\tilde{\xi}_\mathbf{k}(i\omega_n)]^2 - \tilde{\xi}_\mathbf{k}^2 - \Delta_\mathbf{k}^2.
\]

(D9)

This is further simplified for pointlike impurities and \( d \)-wave superconductivity, for which the off-diagonal matrix elements of the self-energy vanish and \( \Delta_\mathbf{k} = \Delta_k \). Because the gap is renormalized, Anderson’s theorem breaks down and the nonmagnetic impurities reduce the order parameter through pairbreaking.

It is common to ignore the chemical potential shift that comes from the real part of the diagonal elements of \( \Sigma(i\omega_n) \). If one makes this assumption, then the dispersion \( \tilde{\xi}_\mathbf{k} \) is also renormalized; that is, \( \tilde{\xi}_\mathbf{k} = \xi_\mathbf{k} \). We do not make this assumption here because we wish to include the chemical potential shift that comes from impurity doping. We have checked that the shifts in electron density so obtained are the same as in the BdG calculations.

With these approximations, the diamagnetic kernel is

\[
K_{xx}^\text{dia} = \frac{1}{N\beta} \sum_{\mathbf{k},n} \frac{\partial^2 \tilde{\xi}_\mathbf{k}}{\partial k_s^2} \text{Tr}[G_{\mathbf{k}}(i\omega_n) \tau_3]
\]

(D10)

and the paramagnetic kernel is

\[
K_{xx}^\text{para} = \frac{1}{N\beta} \sum_{\mathbf{k},n} \left( \frac{\partial \tilde{\xi}_\mathbf{k}}{\partial k_s} \right)^2 \text{Tr}[G_{\mathbf{k}}(i\omega_n)G_{\mathbf{k}}(i\omega_n)].
\]

(D11)

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