Thermodynamic transitions in inhomogeneous \textit{d}-wave superconductors

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We study the spectral and thermodynamic properties of inhomogeneous \textit{d}-wave superconductors within a model where the inhomogeneity originates from atomic scale pair disorder. This assumption has been shown to be consistent with the small charge and large gap modulations observed by scanning tunnelling spectroscopy (STS) on Bi\textsubscript{2}Sr\textsubscript{2}CaCu\textsubscript{2}O\textsubscript{8+x}. Here we calculate the specific heat within the same model, and show that it gives a semi-quantitative description of the transition width in this material. This model therefore provides a consistent picture of both surface sensitive spectroscopy and bulk thermodynamic properties.

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\textbf{Introduction.} There is accumulating evidence from STS that at least some of the families of superconducting cuprate materials are “intrinsically” inhomogeneous at the nanoscale\textsuperscript{1,2,3,4}, in the sense that all samples of the given material exhibit electronic disorder over length scales of $\sim 25\text{Å}$ which cannot be removed by any standard annealing procedure. In particular, differential conductance maps of Bi\textsubscript{2}Sr\textsubscript{2}CaCu\textsubscript{2}O\textsubscript{8+δ} (BSCCO) reveal the existence of spectral peaks reminiscent of the coherence peaks of a homogeneous \textit{d}-wave superconductor whose energy varies by a factor of 2-3 over the sample. There is currently widespread interest in this phenomenon, but no consensus as to its origin, nor even as to whether the gaps between these peaks may be taken to correspond directly to the local superconducting order parameter in the sample, or whether some are related to a second, competing order\textsuperscript{5}. While these experiments have been proposed as evidence of a general nanoscale inhomogeneity in cuprates, there have been several criticisms. One class of objections points out that other materials, particularly YBa\textsubscript{2}Cu\textsubscript{3}O\textsubscript{7−δ} (YBCO) exhibit much narrower NMR linewidths, suggesting that in this material at least, the electronic inhomogeneity cannot be a pronounced property of the bulk\textsuperscript{6}. A stronger critique has been offered by Loram \textit{et al.}\textsuperscript{7} even with regard to the BSCCO material. These authors point out that if one associates with each “gap patch” a local doping corresponding to the hole density required to produce a gap of that size in the average phase diagram, then the distribution of the doping would be of order $\Delta p \sim 0.1$. Such a large range of local doping levels would in turn produce a large inhomogeneous distribution of regions with different $T_c$’s, sufficient to yield a transition width $\Delta T_c$ of order the average $T_c$ itself. Since this is contrary to specific heat measurements\textsuperscript{2,3}, these authors conclude that the inhomogeneity is either a surface phenomenon, or represents a distribution of antinodal scattering lifetimes rather than of order parameters. Clearly it is very important to determine whether results from surface sensitive probes reflect the bulk properties of these materials.

Recently, localized resonances were imaged by STS at a bias of $-960$ meV and identified as the O interstitials primarily responsible for doping the BSCCO material\textsuperscript{10}. This work also noted the strong positive correlation between the positions of these dopants and the local spectral gap, and furthermore argued that the charge inhomogeneity in the sample was considerably smaller than anticipated on the basis of previous scanning tunnelling topographs integrated to smaller bias voltages. Nunner \textit{et al.}\textsuperscript{11} then argued that these and a number of other experimental observations\textsuperscript{12} could be explained by the simple hypothesis that the disorder caused by the dopant atoms was primarily in the Cooper rather than density channel, i.e. that each dopant modulated the BCS pair interaction on an atomic scale.

In this Letter we argue that the lack of strong charge modulations within the pair disorder model of Ref.\textsuperscript{11} also allows one to avoid the argument of Loram \textit{et al.}\textsuperscript{7} regarding the transition width. The model, with parameters chosen to reproduce the gap maps and other correlations of the STS data on BSCCO, is shown to yield results within mean field theory which are consistent with a relative specific heat transition width, $\Delta T_c/T_c$, of $\sim 20\%$, despite the fact that at $T = 0$ gap modulations of order $100\%$ are observed. The observed widths must therefore be attributable to pair disorder alone since the samples are homogeneous at the mesoscale (as determined, e.g., by optical and scanning electron microscopy). Predictions of the theory for the formation and persistence of superconducting islands near the transition should be verifiable by high-temperature STS measurements.

\textbf{Model.} There are many studies of inhomogeneous superconductivity\textsuperscript{12}, only few of which are directly relevant to the questions addressed below. If one adds nonmagnetic disorder to an ordinary \textit{s}-wave superconductor, one expects only small changes in superconducting one-particle properties such as the gap and $T_c$ due to Anderson’s argument\textsuperscript{14}. Nevertheless, as disorder increases and the mean free path $\ell$ becomes comparable to the Fermi wavelength $\lambda_F$, localization effects can destroy
superconductivity. Recently, local aspects of this transition were studied by Ghosal et al. [15], who showed using numerical solutions of the Bogoliubov-de Gennes (BdG) equations including nonmagnetic disorder that near the transition the highly disordered system separates into islands with finite order parameter surrounded by an insulating sea. In the d-wave case, where ordinary disorder is pair breaking and the transition occurs when \( t' \approx \xi_0 \), where \( \xi_0 \) is the coherence length, several numerical studies have investigated local properties of disordered systems [16, 17, 18, 19]. Only Ref. 16 considered the finite temperature transition, however, pointing out that the inhomogeneity induced by ordinary disorder (in the finite temperature transition, however, pointing out considerable higher than the \( \xi_0 \) predicted by the usual disorder-averaged theory of \( T_c \) suppression, analogous to the theory of Abrikosov and Gorkov [20]). The width of the transition was not discussed explicitly in this work, nor was the effect of pairing disorder considered.

More recently, it has been proposed that \( T_c \) can even be increased by local inhomogeneities compared to its value for the homogeneous system [22, 23]. Within a weak coupling BCS framework it was shown that periodic modulations of the pairing and/or electron density can lead to a substantial enhancement of \( T_c \), when the characteristic modulation length scale is of order \( \xi_0 \) [22]. This agrees with recent numerical calculations for the attractive Hubbard model with various disorder distributions of the interaction, verifying that \( T_c \) enhancement by inhomogeneity is a general result arising from the proximity effect [23]. These calculations did not investigate the details of the transition below the \( T_c \), where the local order parameter \( \Delta \) first nucleates.

In the following model calculation we use the standard d-wave BCS Hamiltonian

\[
\hat{H} = \sum_{(ij)\sigma} t_{ij} \hat{c}_{i\sigma}^\dagger \hat{c}_{j\sigma} + \sum_{i\sigma} (V_i - \mu) \hat{c}_{i\sigma}^\dagger \hat{c}_{i\sigma} + \sum_{(ij)} \left( \Delta_{ij} \hat{c}_{i\uparrow}^\dagger \hat{c}_{j\downarrow} + \text{H.c.} \right),
\]

where in the first term we include nearest \( t \) and next-nearest \( t' = -0.3t \) neighbor hopping. For the chemical potential \( \mu \), we set \( \mu = -t \) in order to model the Fermi surface of BSCCO near optimal doping \( \sim 16\% \). \( \sum_{(ij)} \) denotes summation over neighboring lattice sites and \( j \). Disorder is included by the impurity potential \( V_i = V_0 f_i \) where \( f_i = \sum_s \exp(-r_{is}/\lambda)/r_{is}, \ r_{is} \) being the distance from a defect \( s \) to the lattice site \( i \) in the plane. Distances are measured in units of \( a \), where \( a \) is the Cu-Cu distance. Note that the particular Yukawa form of \( f_i \) is merely a convenient way to vary the smoothness of the potential landscape through the parameter \( \lambda \). The d-wave order parameter \( \Delta_{ij} = g_{ij} (\hat{c}_{i\uparrow} \hat{c}_{j\downarrow} - \hat{c}_{i\downarrow} \hat{c}_{j\uparrow})/2 \) is determined self-consistently by iterations of

\[
\Delta_{ij} = \frac{g_{ij}}{2} \sum_n \left( u_n(i) v_n(j) + v_n(i) u_n(j) \right) \tanh \left( \frac{E_n}{2T} \right),
\]

until convergence is achieved. Here, \( \{ E_n, u_n, v_n \} \) is the eigensystem resulting from diagonalization of the BdG equations associated with Eq. (1). The pairing interaction \( g_{ij} \) varies in space relative to its homogeneous value \( g_0 = 1.16t \) as \( g_{ij} = g_0 + \delta g (f_i + f_j)/2 \), where \( \delta g \) is the modulation amplitude and \( i, j \) are nearest neighbors. In the following, when including potential \( (\tau_3 \text{ channel}, V_0 \neq 0) \) or pair \( (\tau_1 \text{ channel, } \delta g \neq 0) \) disorder, we make sure to adjust \( \mu \) and \( g_0 \), respectively, in order to maintain the same doping and average gap as the corresponding homogeneous system. Note that in this approach the inclusion of spatial pair potential variations is purely phenomenological. However, a likely candidate for the pairing modulations is the oxygen dopants. Various possible microscopic origins for this unusual disordered state have been discussed in Refs. [11, 12]. There it was also argued that \( \tau_3 \) disorder alone cannot account for the STM data.

**Results.** Fig. 1(a) shows, in a field of view (FOV) of 49nm × 49nm, the experimental LDOS at -960 meV (a), and the corresponding gap map (b) obtained by McElroy et al. [10]. The bright resonances in (a) reveal the location of the oxygen dopants. In Fig. 1(c) we show the impurity potential generated by using each of these dopants (833 in the above FOV) as a defect located out of the CuO₂ plane. This map resembles (a) as it should. The experimental FOV is modeled as a 90 × 90 lattice system rotated 45 degrees with respect to the 3.83 Å long Cu-Cu bonds, i.e. it includes 2 × 90 × 90 sites and is aligned with the experimental FOV [24]. The theoretical

![Figure 1](attachment:figure1.png)

**FIG. 1:** (Color online.) (a) Experimental \( dI/dV \) map [arb. units] at -960 meV of an optimally doped BSCCO sample from Ref. [10]. (b) Experimental gap map [meV] in the same region as (a). (c) The theoretical impurity potential extracted from (a) assuming distance from plane \( r_z = 0.5 \) and \( \lambda = 0.5 \). (d) The gap map resulting from using (c) as the ‘pairing potential’ in the BdG equations with \( \delta g = 1.5t \). Both (c) and (d) are shown in units of \( t \).
gap map (as extracted from the LDOS) using (c) as a $\tau_1$ potential with $\delta g = 1.5t$ at $T = 0$ is shown Fig. 1(d). We find that gap maps consistent with experiment are found for $1.0 < \delta g/t < 2.0$. The correlation coefficient between the gap maps (b) and (d) is $0.17^{[26]}$, a reasonably large value given the simplicity of the pure $\tau_1$ calculation and the fact that the correlation between the dopant positions and the experimental gap map is $0.3^{[11]}$.

Now we turn to the discussion of the results for $T \neq 0$. Below, we set $N = 50$ (corresponding to the upper left 5/9th square of each image in Fig.1) and study the resulting order parameter (OP) maps, entropy $S(T)$, and specific heat $C(T)$ with focus on the temperature region near $T_c$. The number of iterations necessary for converged results increases dramatically near $T_c$, whereas just a few degrees away from $T_c$ we typically find that 25-50 iterations suffice. In Fig. 2 we show the OP map $\Delta_1 = (\Delta_{i,+x} + \Delta_{i,-x} - \Delta_{i,+y} - \Delta_{i,-y})/4$, for temperatures near $T_c$. The bulk transition temperature for the homogeneous system is $T_{c0} = 0.182t$. Here, one clearly sees how separate islands of finite $\Delta_1$ start forming above $T_{c0}$ when cooling down and eventually overlap at lower $T$. In principle, STS measurements at $T$ close to $T_c$ should be able to probe these superconducting islands.

In order to address the effect of the inhomogeneity on the transition widths, we first calculate the quasiparticle entropy $S(T)$ according to the well-known expression

$$S = -2 \sum_{E_n > 0} [f(E_n) \ln f(E_n) + f(-E_n) \ln f(-E_n)],$$

where $f(E)$ is the Fermi distribution function. From the resulting entropy $S(T)$ curve we extract the specific heat at constant volume $C(T)$ by the usual expression $C(T) = T(\partial S/\partial T)$. Fig. 3(a) compares the electronic specific heat for the clean and pair disordered systems described by different disorder strengths $\delta g$. In Fig. 3(b) we compare $C(T)$ for the different Nambu channels $\tau_1$ and $\tau_3$. We can check the BdG results by comparing to the pure "infinite" system result (black, dashed line in Fig. 3(a)) obtained by replacing $E_n \rightarrow E_k = \sqrt{k_x^2 + \Delta_k^2}$, with $\Delta_k = \Delta_0(T)(\cos k_x - \cos k_y)/2$ with $\Delta_0(T)$ solved selfconsistently from the usual gap equation. It is clear that the $50 \times 50$ BdG lattice problem is large enough to capture the sharp transition width of the clean system.

Conventional potential disorder corresponding to $\sim 10\%$ dopant atoms causes only negligible broadening in the specific heat transition width compared to the pair disordered case. This is shown in Fig. 3(b), where both curves are produced from the impurity distribution in Fig. 1(c). The transition is sharp because potential scatterers cause local suppressions of the order parameter, whereas in interstitial regions $\Delta$ decreases with temperature in a manner similar to the pure system. We have also studied distributions of strong scatterers at the percent level, and find similarly sharp transitions. On the other hand, potential disorder smooth on a scale of $\xi_0$ can lead to large modulations of $\mu$ which, even within BCS theory, will give larger transition widths. In this case, however, the associated local spectra are not consistent with STS measurements, as shown in Ref. 11.

As evident from 3(a), pairing disorder with parameters fixed to yield reasonable variations of the gap size and graininess at low temperature[11] leads to a broadened transition width similar to the experimental observations[2]. This is shown more clearly in Fig. 4 where we compare the experimental specific heat $C(T)$.
for clean YBCO and optimally doped BSCCO near $T_c$, 6 with our results for the clean and pair disordered case. It is clear that in the case of YBCO, the transition is extremely narrow, and the deviations from the mean-field treatment discussed here are consistent with weak 3DXY critical fluctuations over a range of a few degrees.27, 28 On the other hand, in the case of BSCCO, it has been clear for some time that the 3DXY description of the specific heat transition is not appropriate, despite the fact that according to a naive application of the Ginzburg criterion 3DXY fluctuations should be more visible,9 and it has in fact been discussed as closer to a Bose-Einstein condensate (BEC) with specific heat exponent of $\alpha = -1$, although measured values are closer to $\alpha = -0.7$.3 This interpretation remains controversial, however. Other authors have suggested that a 3DXY divergence in $C(T)$ cut off by bulk nanoscale inhomogeneities over a small range $\Delta T_c/T_c \sim 0.03$ could be consistent with the data.7, 29 Here we have put forward a similar scenario, but our microscopic model implies that the inhomogeneities dominate over a larger range $\Delta T_c/T_c \sim 0.1 - 0.2$ to be consistent with STS. Additional contributions from critical fluctuations must be added to the mean field effects discussed here to obtain a complete description.

Conclusions. We have presented theoretical calculations for $d$-wave superconductors with atomic scale pair disorder, using impurity parameters appropriate to reproduce semi-quantitatively the gap maps produced by STM experiments on optimally doped BSCCO, and shown that the experimental specific heat transition in this system can be explained by this model as well. This suggests that substantial nanoscale electronic inhomogeneity is characteristic of the bulk BSCCO system.

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