Properties of the phonon-induced pairing interaction in YBa$_2$Cu$_3$O$_7$ within the local density approximation

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The properties of the phonon-induced interaction between electrons are studied using the local density approximation (LDA). Restricting the electron momenta to the Fermi surface we find generally that this interaction has a pronounced peak for large momentum transfers and that the interband contributions between bonding and antibonding band are of the same magnitude as the intraband ones. Results are given for various symmetry averages of this interaction over the Fermi surface. In particular, we find that the dimensionless coupling constant in the $d$-wave channel $\lambda^d$, relevant for superconductivity, is only 0.022, i.e., even about ten times smaller than the small value of the $s$-wave channel. Similarly, the LDA contribution to the resistivity is about a factor 10 times smaller than the observed resistivity suggesting that phonons are not the important low-energy excitations in high-$T_c$ oxides.

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INTRODUCTION

The relevance of phonons for the low-energy properties of high-$T_c$ oxides is presently rather controversially discussed. Some experiments suggest that the charge carriers near the Fermi surface interact only weakly with phonons. Examples are the magnitude and temperature dependence of the resistivity [1] and the rather small effects in the phonon spectrum caused in general by superconductivity near or below the transition temperature $T_c$ [2]. For instance, the superconductivity-induced softening and width of the zone-center buckling mode in YBa$_2$Cu$_3$O$_7$ corresponds to a rather small electron-phonon (ep) coupling constant [3]. The recently measured isotope dependence of angle-resolved photoemission spectra (ARPES) [4] do not indicate any isotope dependence of the total electronic band width for binding energies down to 200 meV. This shows the inapplicability of a simple polaron picture, caused by a strong ep interaction, to cuprates because the associated band narrowing would be sensitive to different isotopes [5].

On the other hand, there exist many observations which have been taken as evidence for a substantial ep interaction in the cuprates. The measured isotope effects on the transition temperature $T_c$ [4] and the the superfluid density $\lambda$ [6], especially in underdoped samples, are two examples. Another example is the large width and softening observed in bond-stretching phonons in hole-doped cuprates in a very small region in $k$-space [7]. Similarly, the absence of a quasi-particle peak in strongly underdoped Ca$_{2-x}$Na$_x$CuO$_2$Cl$_2$ [8] as well as the doping dependence of a magnetic transition in the frequency-dependent conductivity were attributed to phonons [10, 11]. Finally, electronic self-energy effects in the phonon energy region have been observed by ARPES [12] and by scanning tunneling microscopy [13] and interpreted in terms of a coupling of electrons to a bosonic mode. Whether this mode is related to phonons is presently unclear but if this is the case the ep coupling and phonons would certainly be important for the low-energy physics of cuprates.

Unfortunately, there exists for virtually every interpretation of experiments in favor or disfavor of phonons alternative explanations. For instance, the large isotope effects on $T_c$ observed in the underdoped region do not indicate necessarily a large ep coupling but may be caused by the pseudogap [14]. Similarly, the bosons which interact with electrons near the Fermi surface in the interpretation of ARPES data may be not phonons but, for instance, spin fluctuations [15, 16, 17]. In view of these uncertainties it seems useful to investigate the properties of the ep coupling independently from any interpretation of experiments, i.e., from first-principles using the local density approximation (LDA). To this end we extend recent investigations on phonon-induced electronic self-energy effects [18, 19] and study the momentum, frequency dependence and magnitude of the phonon-induced interaction between electrons in detail. Though our approach is based on the LDA and deals only with the stoichiometric case it takes many features of these systems such as the complicated phonon spectrum and screening properties self-consistently and realistically into account.

COMPUTATION OF THE PHONON-INDUCED INTERACTION

The retarded, phonon-induced electron-electron interaction, multiplied by -1 for convenience, is given in mo-
The amplitudes $g_a$ for the electronic momentum space by $21$, 

$$ V(k\nu, k + q\mu, \omega) = \sum_j |g_j(k\nu, k + q\mu)|^2 \frac{\omega_{qj}}{\omega_{qj} - (\omega + im)^2}. $$

$g_j(k\nu, k + q\mu)$ denotes the renormalized amplitude for a transition from the electronic state with momentum $k$ and band index $\nu$ to the state with momentum $k + q$ and band index $\mu$ creating (annihilating) a phonon with branch label $j$ and momentum $q (-q)$. $\omega_{qj}$ denote the phonon frequencies and $m$ a positive infinitesimal. The amplitudes $g_j$ are obtained within the LDA with the efficient linear-response technique $21, 22$. We used a 36x36x4-mesh for the electronic momentum $k$ within the Brillouin-zone of YBa$_2$Cu$_3$O$_7$, while phonon frequencies and the self-consistent electron-phonon potential were calculated for transferred momenta $q$ on a coarser 12x12x4-mesh. Because even for this reduced mesh an exact calculation was numerically very demanding, we adopted a two-step procedure. First, phonon related quantities were calculated exactly by linear response on a 4x4x2-mesh. Details can be found in Refs. $23$ and $18$. These quantities were then approximated on the 12x12x4-mesh by a Fourier-interpolation technique.

Under the usual assumptions it is sufficient to restrict the electronic momenta $k$ and $k + q$ in $V$ to the Fermi surface. Our momentum meshes allow to put $k$ practically right on the Fermi surface whereas $k + q$ was chosen as near as possible to the Fermi surface. Fermi surface averages were calculated including also a Gaussian for the one-particle energies with width $\delta$. The Fermi surface in YBa$_2$Cu$_3$O$_7$ consists essentially of three bands $\nu, \mu = A, B, C$ where $A, B,$ and $C$ denote the antibonding, bonding and chain band, respectively.

RESULTS

Momentum dependence

The upper panels of Fig. 1 show the intraband contribution $\nu = \mu = B$ to the static interaction $V(k\nu, k'\mu, 0)$. We used $k_z = 0.125$ and $q_z = 0$, i.e., all electronic momenta are restricted to the plane $k_z = 0.125$. The left part in this figure refers to a fixed momentum $k$ at the nodal, the right part at the antinodal Fermi point. Symbols indicate electronic states whose energies differ from the Fermi energy by less than a certain threshold energy. The large filled circles refer to 0.1 eV, the small filled circles to 0.2 eV, the open circles to 0.3 eV. Lines approximate the Fermi surface and are obtained by a linear interpolation between neighboring mesh points. The position in $k$-space of the mesh points corresponding to the different circles is shown in the insets together with the Fermi line depicted by a solid line. $V$ is presented as a function of the angle $\alpha$ at $S = (\pi, \pi)$ between the vectors $(k_x' - \pi, k_y' - \pi)$ and $(-\pi, 0)$. Varying $\alpha$ between 0 and $2\pi$ means that $k'$, seen from the point $S$, moves around the Fermi surface in the anticlockwise sense starting from the antinodal point $\sim (0, \pi)$. The angle $\alpha$ is illustrated in the left inset. It is rather straightforward to construct a continuous curve for $V$ from the discrete points discarding only a very few points which correspond to momenta $k'$ rather far away from the Fermi surface and thus should be omitted.

The right potential curve in Fig. 1 should be symmetric with respect to $\alpha = \pi$ due to the orthorhombic symmetry which is roughly fulfilled for our discrete mesh. If tetragonal symmetry would apply the left potential curve should be symmetric with respect to $\alpha = 5\pi/4$ which holds approximately. Finally, if $V$ depends only on the transferred momentum $q$, the left potential curve, shifted rigidly by the angle $\alpha = -\pi/4$, would coincide with the right potential curve. This is qualitatively the case, for instance, the two dominating maxima are close to each other after such a shift. Quantitatively there are, however, differences, for instance, the heights of the maxima differ by about 30% and the (small) values at zero momentum transfer by about a factor 3 after the shift. Remarkable is the large variation of $V$ by about a factor of 5 between small and large momentum transfers, i.e., between $\alpha = 0$ and $\pi$ in the case of the right potential curve. Interesting for the relation between state and transport relaxation times is that $V$ is larger for large than for small momentum transfers.

The lower panels in Fig. 1 show potential curves for interband scattering between the antibonding and bond-
near the Fermi surface described by the solid lines. Similar as in Fig. 1. Insets show the considered mesh points $k$.

![FIG. 2: Phonon-induced interaction $V(kA, k'A, 0)$ (upper part) and $V(kB, k'A, 0)$ (lower part) for a fixed momentum $k$ at the nodal (left diagrams) and antinodal (right diagrams) Fermi point as a function of $k'$, encircling the Fermi surface similar as in Fig.1. Insets show the considered mesh points near the Fermi surface described by the solid lines.](image)

The interband contribution between the bonding and antibonding band, shown in the lower part of Fig. 2, looks qualitatively similar as the interband term between the antibonding and the bonding band of Fig. 1 exhibiting well-pronounced maxima at large momentum transfers. At small momentum transfers electrons near the nodal direction interact only weakly whereas those in the antinodal direction develop a second and rather sharp peak in $V$ at $\alpha = 0$ which, to a lesser degree, was also present in the intraband contribution of the antibonding band. The absolute magnitude of $V$ is in all considered cases similar which means that interband and intraband terms are comparable in magnitude.

Fig. 3 shows potential curves for interband scattering between the band $A$ and $B$, respectively, and the chain band $C$. Due to the geometry of the chain band the angle $\alpha$ is restricted to some region around $\pi/2$ and $3\pi/2$. The potentials peak in general around these to values but the absolute values are rather small compared, for instance, to those in Fig. 1.

![FIG. 3: Phonon-induced interaction $V(kB, k'C, 0)$ (upper part) and $V(kA, k'C, 0)$ (lower part) for a fixed momentum $k$ at the nodal (left diagrams) and antinodal (right diagrams) Fermi point as a function of $k'$, encircling the Fermi surface similar as in Fig.1. Insets show the considered mesh points near the Fermi surface described by the solid lines.](image)

The dimensionless coupling function $\lambda_{\nu}(k)$ [20] can be written in terms of $V$ as,

$$\lambda_{\nu}(k) = 2 \sum_{k', \mu} V(k\nu, k'\mu, 0) \delta(\epsilon_{k'\mu}).$$  \hspace{1cm} (2)

Fig. 4 shows $\lambda_{\nu}(k)$ for $\nu = A$ (red squares joined by straight lines), $\nu = B$ (black circles) and $\nu = C$ (blue triangles) for $\delta = 0.2$ eV as a function of the angle $\alpha$. Using $k_\pi = 0.125$ and $k_\gamma = 0.375$ yields two curves denoted by filled and empty symbols, respectively, which are close to each other illustrating the weak dependence of $\lambda(k)$ on $k_\gamma$. Both for the bonding and antibonding bands $\lambda(k)$ is approximately symmetric with respect to $\alpha = \pi/4$ reflecting the tetragonal symmetry of isolated layers. Moreover, the coupling functions are smaller along the nodal direction by about 20 to 30% compared to the antinodal direction which characterizes the anisotropy of $\lambda(k)$ in the layers. The Fermi surface of the chain band starts in our plot only somewhat below the nodal direction and yields a rather strong and rapidly varying coupling function $\lambda_C(k)$ reflecting the strong perturbation of tetragonal symmetry by the chains.

Dimensionless coupling constants $\lambda_{\nu}^\alpha$ with $\alpha = \ldots$
by Gaussians with width $\delta$ band $\nu$ (empty symbols). Angle $k$ (blue triangles) for according to Eq. (2) those for the remaining “non-trivial” symmetries. Acquisition of coupling constants for the three bands is similar in total coupling constant relevant for superconductivity.

The numbers in the columns 2-4 of Table I show that high-frequency part of transitions. Considering $\delta = 0.1$ eV instead of 0.2 eV does not change much our results: For instance, the first three numbers in the first line become 0.226, 0.229, 0.230, respectively. This suggests that our momentum nets are adequate for calculating average quantities such as $\lambda^\alpha$.

The most interesting numbers in the table are those in the last column which determine the phonon contribution to superconductivity. Isotropic $s$-wave dominates by far but this component is ineffective because of the strong Coulomb repulsion. In all other symmetry channels $\lambda$ is very small. This holds in particular for the $d$-wave channel where it is only 0.022, i.e., one order of magnitude smaller than the isotropic $s$-wave value. This value is rather stable with respect to $\delta$. For instance, it changes from 0.022 to 0.027 if $\delta = 0.2$ eV is reduced to $\delta = 0.1$ eV. Keeping only intraband contributions one obtains 0.021, i.e., the interband transitions are negligible in this case.

Using the Kramers-Kronig transformation for $V$ in Eq. (3) $\lambda^\nu$ can be written as an integral from zero to infinity over a frequency-dependent coupling function $\lambda^\nu(\omega)$.

![FIG. 4](image-url) (Color online) Coupling functions $\lambda_A(k)$ (red squares joined by straight lines), $\lambda_B(k)$ (black circles), and $\lambda_C(k)$ (blue triangles) for $k_z = 0.125$ (filled symbols) and $k_z = 0.375$ (empty symbols). Angle $\alpha$ is the same as in Figs. 1, 2, and 3.

![FIG. 5](image-url) (Color online) Coupling functions $\lambda^s(\omega)$ (black dashed line) and $\lambda^d(\omega)$ (red solid line) as a function of frequency using a width of 1 meV for the phonon modes.

The numbers in the columns 2-4 of Table I show that the coupling constants for the three bands are similar in magnitude for each symmetry component. The isotropic $s$-wave component is in all cases substantially larger than those for the remaining “non-trivial” symmetries. According to Eq. (2) $\lambda_\nu(k)$ and thus also the above coupling constants contain both intraband ($\mu = \nu$) and interband ($\mu \neq \nu$) contributions. Keeping only the intraband parts diminishes substantially the numbers in the table. For instance, for the isotropic $s$-wave channel (first line) the numbers for $A, B, C$ change to 0.090, 0.069, 0.074, respectively. This means that interband transitions contribute much more to the coupling functions than the intraband transitions.

In the case of a momentum-independent $V$ where it is only 0.022, i.e., one order of magnitude smaller than the isotropic $s$-wave value. This value is rather stable with respect to $\delta$. For instance, it changes from 0.022 to 0.027 if $\delta = 0.2$ eV is reduced to $\delta = 0.1$ eV. Keeping only intraband contributions one obtains 0.021, i.e., the interband transitions are negligible in this case.
to the buckling and breathing phonon frequencies, respectively. However, these peaks have different signs and thus cancel each other to a large extent in $\lambda_{tr}$. We note that the positive peak in $\lambda_t^/(\omega)$ at 27 meV is related to an oxygen buckling-type vibration with odd symmetry with respect to the CuO$_2$ bilayer.

In Fig. 6 we show a comparison between the frequency dependence of the total coupling functions with $s$ and $s'$ (upper panel) and with $p_x$ and $p_y$ (lower panel) symmetries. As expected the $s$-wave coupling function is much larger throughout the whole frequency interval compared to the other symmetries. Remarkable is that the couplings with $p_x$ and $p_y$ symmetries are mainly negative over the whole frequency region and that they differ from each other reflecting the presence of the chain band and the broken tetragonal symmetry.

**Resistivity**

The resistivity in $i$-direction is determined by the transport coupling constant $\lambda_{tr}^i$, given by [20]

$$\lambda_{tr}^i = 2 \sum_{k, k', \nu, \mu} V(k\nu, k'\mu, 0) \delta(\epsilon_{k\nu}) \delta(\epsilon_{k'\mu}) \cdot$$

$$\left( \langle v_{i}^2(k\nu) \rangle - \langle v_{i}(k\nu) \rangle \langle v_{i}(k'\mu) \rangle \right) / \langle v_{i}^2 \rangle,$$  \hspace{1cm} (4)

where $\langle v_{i}^2 \rangle$ denotes the average of $v_{i}^2(k\nu)$ over all pieces of the Fermi surface. Our calculation yields $\lambda_{tr}^{x} = 0.256$, $\lambda_{tr}^{y} = 0.272$, and $\lambda_{tr}^{z} = 0.228$. These values are close to $\lambda_t$ illustrating the fact that the pronounced momentum dependence of $V$ in Figs. 13-15 not necessarily reflects itself in the above coupling constants. Generalizing $V$ in Eq. 34 to a finite frequency and taking the imaginary part the right-hand side of this equation is proportional to the function $\alpha_{tr}^i F(\omega)$ from which the temperature dependence of the resistivity can be calculated using Eqs. (4) and (7) of Ref. [24]. The results are shown in Fig. 7.

**FIG. 7: (Color online) Temperature dependence of the resistivity $\rho$, along the direction $i$.**

With increasing temperature the curves reach between 120-150 K a quasi-linear regime which approaches in a very smooth way the true linear region above the highest phonon frequency at around 1000K (not shown in Fig. 7). A typical experimental value is $\rho_x(300K) = 290 \cdot 10^{-6}$ Ohm cm [1], i.e., only about 10 % of the experimental scattering is due to phonons. Also the noticeable deviations from a linear law between $T_c = 90K$ and about 150 K in the theoretical curves seems to be in conflict with the experiment for optimally doped samples.

**DISCUSSION**

Our approach treats electronic correlations only in an approximate way and its applicability to real YBa$_2$Cu$_3$O$_7$ may be questioned. Unfortunately, the calculation of corrections to our results due to strong electronic correlations is difficult and presently not very reliable. However, it seems very improbable that our main results, namely that $\lambda_t$ and $\lambda_{tr}$ are about a factor 50 and 10 too small to account for $T_c$ and $\rho$, respectively, will be substantially changed by correlations. Experimental data on the width of some phonons are interesting in this respect. The anomalous broadening of the buckling phonon at the zone center has been quantitatively explained with LDA results for the ep-interaction, once a large anharmonic contribution was subtracted [3]. On the other hand, bond-stretching phonons at low temperatures show anomalies around the wave vector $(0, 0.25, 0)$ in form of a sharp and localized softening and large widths $\lambda_{tr}$ which clearly are beyond the LDA [25]. At higher temperatures, where these anomalies are not present, the measured widths are still sizable. Assuming that these widths are entirely due to the ep-interaction our LDA calculation could only account for about 20 % of these widths. However, part of the widths could also be caused by anharmonicity like in Ref. [3]. Furthermore, it is not clear how much these rather localized anomalies
in k-space contribute to Fermi surface averaged quantities like $\lambda^{tr}$. A large increase of $\lambda^{tr}$ due to correlation-enhanced coupling to bond-stretching phonons could account for the missing scattering contribution found in the LDA calculation of $\rho$. On the other hand, its temperature dependence would be in conflict with the observed linear temperature between $T_c$ and room temperature because the onset of the linear dependence in Fig. 7 would be shifted to higher temperatures due to the high frequency of bond-stretching phonons.

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