Extracting the electron–boson spectral function $\alpha^2 F(\omega)$ from infrared and photoemission data using inverse theory

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We present a new method of extracting electron-boson spectral function $\alpha^2 F(\omega)$ from infrared and photoemission data. This procedure is based on inverse theory and will be shown to be superior to previous techniques. Numerical implementation of the algorithm is presented in detail and then used to accurately determine the doping and temperature dependence of the spectral function in several families of high-$T_c$ superconductors. Principal limitations of extracting $\alpha^2 F(\omega)$ from experimental data will be pointed out. We directly compare the IR and ARPES $\alpha^2 F(\omega)$ and discuss the resonance structure in the spectra in terms of existing theoretical models.

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I. INTRODUCTION

The electron–boson spectral function is one of the most important properties of a BCS superconductor. In conventional superconductors the electron-phonon spectral function has been successfully obtained using tunneling and infrared (IR) spectroscopy. The situation is more complicated in cuprates where the mechanism of superconductivity is still a matter of debate. Based on IR data it was suggested very early that charge carriers in cuprates might be strongly coupled to some collective boson mode. It was subsequently proposed that this collective mode might be magnetic in origin. Within this scenario electrons are strongly coupled to a so called “41 meV” resonance peak observed in INS (Refs. [11]). The peak is believed to originate from antiferromagnetic spin fluctuations that persist in the superconducting state; coupling of electrons to this mode in turn leads to Cooper pairing. However recently this view was challenged by a proposal that charge carriers might be strongly coupled to phonons. This controversial suggestion has revitalized the debate about whether a collective boson mode is responsible for superconductivity in the cuprates. An accurate and reliable determination of the electron–boson spectral function has become essential.

In this paper we propose a new way of extracting the spectral function from IR and Angular Resolved Photoemission Spectroscopy (ARPES) data. The proposed method is based on inverse theory, and will be shown to have numerous advantages over previously employed procedures. An advantage of the method is that it eliminates the need for differentiation of the data, that was previously the most serious problem. The inversion algorithm uncovers extreme sensitivity of the solution to smoothing, and offers a smoothing procedure which eliminates arbitrariness. Since the spectral function is convoluted in the experimental data, some information is inevitably lost; we will use inverse theory to set the limits on useful information that can be extracted from the data. Unlike previous techniques which are valid only at $T=0$ K, the new method can be applied at any temperature.

The paper is organized as follows. First in Section II we outline the numerical procedure of solving integral equations. In Section III we demonstrate the usefulness of the new method by applying it to previously published data for $\text{YBa}_2\text{Cu}_3\text{O}_{7-\delta}$ ($\text{Y123}$). In Section IV model calculations of spectral function will unveil some important problems encountered when solving integral equations. Section V discusses the origin of negative values in the spectral function and methods for dealing with them. In Section VI the effect superconducting energy gap has on the spectral function will be analyzed. In Section VII we study the temperature dependence of the spectral function for optimally doped $\text{Bi}_2\text{Sr}_2\text{CaCu}_2\text{O}_{8+\delta}$ ($\text{Bi2212}$). In Section VIII inverse theory is applied to ARPES data and the spectral function of molybdenum surface Mo(110) and Bi2212 have been studied. Finally, Section IX contains quantitative comparison of the spectral functions of optimally doped Bi2212, extracted from both IR and ARPES data; the observed results are critically compared against existing theoretical models. In Section X we summarize all the major results.

II. NUMERICAL PROCEDURE

The (optical) scattering rate of electrons in the presence of electron-phonon coupling at $T=0$ K is given by famous Allen’s result:

$$\frac{1}{\tau(\omega)} = \frac{2\pi}{\omega} \int_{0}^{\omega} d\Omega (\omega - \Omega) \alpha^2 F(\Omega),$$

where $\alpha^2 F(\omega)$ is the electron-phonon spectral function. The scattering rate $1/\tau(\omega)$ can be obtained from complex optical conductivity $\sigma(\omega) = \sigma_1(\omega) + i\sigma_2(\omega)$:
\[
\frac{1}{\tau(\omega)} = \frac{\omega_p^2}{4\pi} \frac{\sigma_1(\omega)}{\sigma_1(\omega) + \sigma_2(\omega)},
\]  
(2)

where \(\omega_p\) is the conventional plasma frequency. Recently Marsiglio, Startseva and Carbott\textsuperscript{a} have defined a function \(W(\omega)\):

\[
W(\omega) = \frac{1}{2\pi} \frac{d^2}{d\omega^2} \left[ \omega \cdot \frac{1}{\tau(\omega)} \right],
\]  
(3)

which they claim to be \(W(\omega) \approx \alpha^2 F(\omega)\) in the phonon region\textsuperscript{5}. It is easy to show (by substitution, for example) that Allen’s formula Eq. (1) and Eq. (3) are equivalent expressions, provided \(1/\tau(\omega) = 0\). Eq. (3) is frequently used to extract the spectral function in the cuprates from IR data\textsuperscript{9,14,21,22,23,24,25,26}. Obviously this method introduces significant numerical difficulty since the second derivative of the data is needed. The experimental data must be (ambiguously) smoothed “by hand” before Eq. (3) can be applied, otherwise the noise will be amplified by (double) differentiation and will completely dominate the solution. An alternative approach is to fit the scattering rate with polynomials and then perform differentiation analytically\textsuperscript{20,27}. Note also that although Eq. (3) is valid only at \(T = 0\) K, it is frequently applied to higher \(T\), even at room temperature.

Here we propose a new method of extracting the spectral function. It is based on the following formula for the scattering rate at finite temperatures derived by Shulga et al\textsuperscript{28,29}:

\[
\frac{1}{\tau(\omega, T)} = \frac{\pi}{\omega} \int_0^\infty d\Omega \alpha^2 F(\Omega, T) \left[ 2\omega \coth \left( \frac{\omega + \Omega}{2T} \right) - (\omega + \Omega) \coth \left( \frac{\omega + \Omega}{2T} \right) + (\omega - \Omega) \coth \left( \frac{\omega - \Omega}{2T} \right) \right],
\]  
(4)

which in the limit \(T \to 0\) K reduces to Allen’s result Eq. (1) (Ref. 20). Unlike Eq. (1) which has a differential form Eq. (2), there is no such simple expression for Eq. (4). Therefore in order to obtain \(\alpha^2 F(\omega)\) from Eq. (4) one must apply inverse theory\textsuperscript{30}. Like most inverse problems, obtaining the spectral function from the scattering rate data is an ill-posed problem which requires special numerical treatment. The spectral function appears under the integral, an operator which has smoothing properties. That means that some of the information on \(\alpha^2 F(\omega)\) is inevitably lost. Using inverse theory our goal will be to extract as much of useful information as we can, and set the limits on lost information.

Numerically the procedure of solving an integral equation reduces to an optimization problem, i.e. finding the “best” out of all possible solutions\textsuperscript{30}. Different criteria can be adopted for the “best” solution, such as: 1) closeness to the data in the least square sense (we will call this solution “exact”) or 2) smoothness of the solution. The most useful solution is often a trade-off between these two.

Eq. (4) is a Fredholm integral equation of the first kind\textsuperscript{30} it may be rewritten as:

\[
\frac{1}{\tau(\omega, T)} = \int_0^\infty d\Omega \alpha^2 F(\Omega, T) K(\omega, \Omega, T)
\]  
(5)

where \(1/\tau(\omega, T)\) is experimental data (from Eq. (2)), \(K(\omega, \Omega, T)\) (contains the prefactor \(\pi/\omega\) from Eq. (1)) is a so-called kernel of integral equation, and \(\alpha^2 F(\omega, T)\) is the unknown function to be determined. When discretized in both \(\omega\) and \(\Omega\) Eq. (4) becomes:

\[
\frac{1}{\tau(\omega_i, T)} = \sum_{j=1}^N \Delta \Omega_j \alpha^2 F(\Omega_j, T) K(\omega_i, \Omega_j, T),
\]  
(6)

with \(i = 1, N\). In matrix form:

\[
\vec{\gamma} = K \vec{a},
\]  
(7)

where vector \(\vec{\gamma}\) corresponds to \(1/\tau(\omega_i, T)\), vector \(\vec{a}\) to \(\alpha^2 F(\Omega_j, T)\) and matrix \(K\) to \(K(\omega_i, \Omega_j, T)\) (Ref. 31). The problem is reduced to finding vector \(\vec{a}\), i.e. the inverse of matrix \(K\). To perform this matrix inversion we adopt a so called singular value decomposition (SVD)\textsuperscript{30} because it allows a physical insight into the inversion process and offers a natural way of smoothing. Matrix \(K\) is decomposed into the following form:

\[
K = U [\text{diag}(w_j)] V^T
\]  
(8)

where \(U\) and \(V\) are orthogonal matrices (\(U^T = U^{-1}\) and \(V^T = V^{-1}\)), and \(\text{diag}(w_j)\) is a diagonal matrix with elements \(w_j\). The inverse of \(K\) is now trivial: \(K^{-1} = V [\text{diag}(1/w_j)] U^T\) and the solution to Eq. (7) is than simply:

\[
\vec{a} = K^{-1} \vec{\gamma} = V [\text{diag}(1/w_j)] U^T \vec{\gamma}.
\]  
(9)

The elements of diagonal matrix \(w_j\) are called singular values (s.v.); they are by definition positive and are usually arranged in decreasing order. If all of them are kept in Eq. (9) the “exact” solution, i.e. the best agreement with the original data, is obtained. If needed, and it almost always is when solving integral equations, the smoothing of the solution (not the experimental data) is achieved by replacing the largest \(1/w_j\) in Eq. (9) with zeros, before performing matrix multiplications. This is a common procedure of filtering out high frequency components in the solution\textsuperscript{30}. 

III. AN EXAMPLE

To demonstrate the usefulness of this procedure we first analyze the existing IR data for underdoped YBa$_2$Cu$_3$O$_{6.6}$ with $T_c=59$ K (Ref. 8). Spectral function $W(\omega)$ for this compound was previously determined using Eq. (3), after $1/\tau(\omega,T)$ had been (heavily) smoothed. Here we apply the numerical procedure described in the previous section on the same data set. We start with $1/\tau(\omega, T=10K)$ data in the range 10-3000 cm$^{-1}$, and form a linear set of 300 equations to be solved (N=300 in Eq. (6)), i.e. 300-element vectors $\vec{a}$ and $\vec{\gamma}$ and a 300x300 matrix $K$ (Eq. (7)). We then decompose matrix $K$ (Eq. (8)) and choose how many of its singular values we are going to keep. Finally we invert the matrix and solve the system for vector $\vec{a}$ (Eq. (9)), i.e. $\alpha^2 F(\omega)$ in the range between 10-3000 cm$^{-1}$, at 300 points.

Left panels of Fig. 1 show results of $\alpha^2 F(\omega)$ calculations for YBa$_2$Cu$_3$O$_{6.6}$ at 10 K, for 6 different levels and/or methods of smoothing. The right panels show the scattering rate $1/\tau(\omega)$, along with the calculated scattering rate $1/\tau_{cal}(\omega)$, obtained by substituting the corresponding $\alpha^2 F(\omega)$ on the left back into Eq. (4). The top panels (A1 and A2) display previously published solutions obtained using Eq. (3) after the data had been smoothed "by hand". The next two panels (B1 and B2) present the "exact" solution using SVD, with all 300 singular values different from zero. This solution does not appear to be very useful (note the vertical scale), although it gives the best agreement between the experimental data $1/\tau(\omega)$ and the right panels the experimental $1/\tau(\omega)$ along with $1/\tau_{cal}(\omega)$ calculated from the corresponding spectral function (Eq. (4)). The top panel shows previously published spectral function obtained from the scattering rate smoothed "by hand". The other five pairs of panels are the data obtained using inverse theory. Different number of singular values are kept in the calculations, which results in different levels of smoothing. Note that the vertical scale in panels B1 and C1 is different.

From Figs. 1 and 2 we conclude that extreme caution is required when performing numerical procedures based on the data smoothed "by hand". In Fig. 2 the strongest peak at around 480 cm$^{-1}$ is fairly robust, although its spectral weight does change a few percent. However the other structures are very dependent on smoothing. The strongest dip shifts from 780 cm$^{-1}$ with 11 s.v. to 760 cm$^{-1}$ with 12 s.v. and 750 cm$^{-1}$ with 13 s.v. In the data smoothed "by hand" it is at 730 cm$^{-1}$. The spectral weight of the dip also varies. It was suggested by Abanov et al. that the main dip, not the peak, is a better measure of the frequency $2\Delta + \omega_s$. However based on our calculations (Fig. 2) the dip is even more sensi-
IV. MODEL CALCULATIONS

The question we must now try to answer is how many s.v. to keep in inversion calculations. To address this issue we have performed calculations based on model spectral function with two Lorentzians:

\[
\alpha^2 F(\omega) = \frac{\omega_p^2\omega^2}{(\omega_a - \omega^2)^2 + (\gamma_a\omega)^2} + \frac{\omega_p^b\omega^2}{(\omega_b^2 - \omega^2)^2 + (\gamma_b\omega)^2},
\]

(10)

with \(\omega_{p,a} = 50,000\text{ cm}^{-1}\), \(\omega_a = 500\text{ cm}^{-1}\), \(\gamma_a = 200\text{ cm}^{-1}\), \(\omega_{p,b} = 250,000\text{ cm}^{-1}\), \(\omega_b = 2,000\text{ cm}^{-1}\) and \(\gamma_b = 800\text{ cm}^{-1}\). This analytic form was chosen to mimic the “real” spectral function in cuprates (see for example Fig. 3 below). From this \(\alpha^2 F(\omega)\) the scattering rate was calculated (not shown) using Eq. (10) and then the formalism of inverse theory (Section II) was applied. Figure 3 shows the model spectral function (gray lines) along with the spectral function determined using inverse theory (black lines). We see that the “exact” solution (with all 300 s.v.) does not agree well with the model; this is due to numerical instabilities induced by smallest s.v. As we reduce the number of s.v. (cut-off the smallest) the agreement improves and for 100 and 50 s.v. the inversion reproduces the original spectral function. As we reduce the number of s.v. further the agreement begins to deteriorate and negative values in \(\alpha^2 F(\omega)\) appear again. Obviously these negative values are not real and simply reflect the fact that too few s.v. do not contain enough information to reproduce the original data. Note however that even with very few s.v. the main features of the spectral function are reproduced, as the main peaks and dips are roughly at correct frequencies (see for example calculations with 20, 15 and 10 s.v.). Their spectral weights are not reproduced though.

The optimal number of s.v. is always a trade–off between numerical precision and closeness to the data. Unfortunately, unlike model calculation shown in Fig. 3 in calculations with real data those two criteria are not well separated. Therefore one must be very careful when quantitatively analyzing the fine structure and their spectral weight in \(\alpha^2 F(\omega)\), as different levels and/or methods of smoothing can cause spurious shifts of the peaks and/or redistribution of their weights. For example visual inspection of \(1/\tau_{\text{cal}}(\omega)\) on the righthand side of Fig. 4 cannot distinguish between different levels of smoothing [compare \(1/\tau_{\text{cal}}(\omega)\) in panels C2, D2 or E2], however even the smallest differences manifest themselves in the spectral functions in the left panels.

An advantage of using inverse theory for extracting \(\alpha^2 F(\omega)\) is that we can quantify the smoothing procedure by specifying the number of s.v. different from zero in Eq. (9), thus eliminating arbitrariness related with smoothing of experimental data “by hand”. This is especially important when quantitatively comparing results from two different \(1/\tau(\omega)\) curves. Note however that if the data sets have different signal-to-noise levels, keeping the same number of s.v. will result in different levels of smoothing. We will encounter this problem below when we study doping dependence of \(\alpha^2 F(\omega)\) in Y123, since available data are from different sources.

Similar problems arise when analyzing temperature dependence of the data. Keeping the same number of singular values is again not the best way to achieve similar levels of smoothing. Fig. 4 shows the absolute values of first (biggest) 200 s.v. at different temperatures. They drop quickly (note the log scale) and such small \(w_i\) produce large oscillations in the solution. To avoid that one cuts–off, i.e. replaces \(1/w_i\) with zeros in Eq. (9). As Fig. 4 shows s.v. are also very temperature dependent, and there are different ways to make the cut. As mentioned above keeping the same number of s.v. different from zero (“vertical cut”) is not a good way, as that would imply including smaller s.v. at higher temperatures and therefore higher frequency components into the solution. In such cases it is better to make “horizontal cuts”, i.e. keep the s.v. in the same range of absolute values. This implies different number of s.v. at different temperatures, but the oscillations in all the solutions should be approximately the same.
FIG. 3: Model calculations of spectral function with two Lorentzians (Eq. 10). The “exact” solution, with all 300 s.v., does not agree well with the model because small singular values produce numerical instabilities in the solution. On the other hand if too few s.v. are kept unphysical negative regions appear. The model spectral function is recovered with 50–100 s.v.

V. PROBLEM OF NEGATIVE VALUES

An obvious problem with these (Figs. 1 and 2) and previous (Ref. 8, 21, 22, 23, 24, 25, 26) calculations is that they all produce non-physical negative values in the spectral function. The latter function is proportional to boson density of states $F(\omega)$ and therefore cannot be negative. The important issue we must address is the origin of these negative values. As shown in Fig. 3 negative values can appear because of numerical problems: either because small s.v. produce numerical instabilities, or because too few s.v. do not contain sufficient information to reproduce the original data. These negative values are not real and can be eliminated either by choosing appropriate number of s.v., or by some other numerical technique, as we will show below.

However, negative values can also have a real physical origin, and they cannot be eliminated by any numerical procedure. Namely, all the methods we have discussed (Eqs. 1, 2 or 3) were developed for normal state, but are frequently used in the (pseudo)gapped state. In order to illustrate the insufficiency of these models to account for a (pseudo)gap in the density of states we have performed inversion calculations on BCS scattering rate. Fig. 5 shows that scattering rate (right panels) calculated within BCS with $\Gamma = 2\Delta = 400 \text{cm}^{-1}$, at $T/T_c = 0.1$. The spectral functions calculated with different number of s.v., i.e. different levels of smoothing are shown in left panels. Surprisingly they look very similar to those produced by coupling of carriers to collective bosonic mode (see Figs. 1 and 2): there is a strong peak roughly at the frequency of the gap, followed by a strong dip and fine structure which is smoothing dependent.

The main issue now is whether one can distinguish between real, physical negative values arising because of the gap in the density of states and those arising because of numerical instabilities. Using inverse theory we can also address this problem. A so called deterministic constraints can be imposed on the solution during the inversion process. These deterministic constraints reduce the set of possible solutions from which the “best” solu-
Fig. 5: Model calculations of spectral function from BCS scattering rate. Left panels display calculated spectral function $\alpha^2 F(\omega)$ and the right panels the BCS scattering rate (also shown with dotted lines in left panels) and $1/\tau_{\text{cal}}(\omega)$ calculated from the spectral function on the left. A gap in the density of states produces similar structure in $\alpha^2 F(\omega)$ as does the coupling to a bosonic mode.

Numerically one applies the constraints during an iterative inversion process. The initial solution $\vec{a}_0$ for the iteration can be obtained either from Eq. (7) or more generally using a so called regularization:

$$K^T \vec{\gamma} = (K^T K + \delta H) \vec{a},$$  \hspace{1cm} (11)

where $H$ is a so called regularization matrix and $\delta$ is a regularization parameter. For $\delta = 0$ (no regularization) Eq. (11) reduces to Eq. (7). Eq. (11) can also be solved using SVD. Once the initial solution $\vec{a}_0$ is found, one applies iteration, imposing the constraint $\alpha^2 F(\omega) \geq 0$ in every step:

$$\vec{a}_{n+1} = P[(I - \beta \delta H) \vec{a}_n + \beta K^T (\vec{\gamma} - H \vec{a}_n)],$$  \hspace{1cm} (12)

where $\beta$ is the iteration parameter and $P$ denotes an operator that sets all the negative values in the solution to zero. The results of these calculations for YBa$_2$Cu$_3$O$_{6.6}$ with $T_c = 59$ K are shown in Fig. 6. The initial solution (top panels) was obtained from SVD with 20 s.v. and no regularization. This solution was then iterated different number of times: 100 (panel B), 200 (C), 500 (D) and 1000 (F). For each intermediate solution the scattering rate $1/\tau_{\text{cal}}(\omega)$ (gray lines) was calculated using Eq. (11).

Clearly as the number of iterations increase the agreement between $1/\tau(\omega)$ and $1/\tau_{\text{cal}}(\omega)$ becomes better, but it never becomes as good as the one with negative values (top panel). It appears that the numerical process converges, although very slowly, to the solution with negative values: some frequency regions in $\alpha^2 F(\omega)$ have simply been cut-off by the program. The position of the main peak is not affected, but its intensity has been reduced significantly. We also emphasize that the structure in the spectral function at $\omega > 1,000 \text{ cm}^{-1}$ is essential for obtaining linear frequency dependence of $1/\tau(\omega)$ up to...
very high frequencies. We will return to this important issue in Section IX below.

Therefore in the case of YBa$_2$Cu$_3$O$_{6.60}$ we have been able to eliminate negative values and at least in principle obtain $\alpha^2 F(\omega)$ which is always positive. This indicates that the structure in the spectral function is (predominantly) due to coupling to bosonic mode and not the gap in the density of states. On the other hand, we have not been able to obtain good BCS scattering rate without negative values in spectral function (not shown). This is not unexpected as the form of spectral function is entirely due to a gap in the density of states (no bosonic mode), which Eqs. (11) and (12) do not take into account. We have encountered similar situation in some cuprates. Fig. 7 displays inversion calculations for several Y123 samples with different doping levels and/or $T_c$: YBa$_2$Cu$_3$O$_{6.60}$ with $T_c = 57$ K (Ref. 34), YBa$_2$Cu$_3$O$_{6.60}$ with $T_c = 59$ K (Ref. 8) and YBa$_2$Cu$_3$O$_{6.95}$ with $T_c = 91$ K (Ref. 34). All calculations are for $T=10$ K, with a fixed number of 15 s.v. As can be seen from Fig. 7 the peak systematically shifts to higher energies as doping and $T_c$ increase: 430 cm$^{-1}$ in $x=6.6$ with $T_c=57$ K, 480 cm$^{-1}$ in the second $x=6.6$ with $T_c=59$ K and 520 cm$^{-1}$ in $x=6.95$. For both 6.6 samples we have been able to obtain relatively good inversions (dashed lines) without negative values in the spectral function. That is not the case for 6.95 sample where without negative values the inversion fails badly (see dashed line in the bottom–right panel). This indicates that the form of scattering rate is probably a combination of coupling to collective mode and a gap in the density of states, as pointed out by Timusk.

VI. ELECTRON-BOSON COUPLING VS. ENERGY GAP

As demonstrated in previous sections similar shapes of $\alpha^2 F(\omega)$ are produced by coupling to bosonic mode and a gap in the density of states when equations for the normal state (Eq. 11) or (12) are used. It is essential to discriminate these two contribution because they usually appear together. To address this problem we have to apply Allen’s formula for the scattering rate in the superconducting state.

$$\frac{1}{\tau(\omega)} = \frac{2\pi}{\omega} \int_{0}^{\omega - 2\Delta} d\Omega (\omega - \Omega) \alpha^2 F(\Omega) \left[ 1 - \frac{4\Delta^2}{(\omega - \Omega)^2} \right]$$

In this equation $E(\omega)$ is the complete elliptic integral of the second kind and $\Delta$ is a gap in the density of states. For $\Delta=0$ Eq. 13 reduces to Eq. 11 for the normal state. Numerically Eq. 13 is again Fredholm integral equation of the second kind and the same numerical procedure for its solution can be used.

We have performed inversion of the data for optimally doped YBa$_2$Cu$_3$O$_{6.95}$ using Eq. 13. Fig. 8 shows inversion calculations for different values of the gap $\Delta$. The top panels display calculations with $\Delta=0$, which is equivalent to previous calculations using Eq. 11 (Fig. 7). As we already discussed it, the spectrum is dominated by a pronounced peak, followed by a large negative deep. Unlike YBa$_2$Cu$_3$O$_{6.6}$ for which this negative deep can, at least in principle, be eliminated, the deep in YBa$_2$Cu$_3$O$_{6.95}$ cannot be eliminated (Fig. 8) and in the previous section we suggested that that is because of the gap. Indeed when finite values of the gap are used in Eq. 13 this negative deep following the main peak is strongly suppressed; calculated spectral function positive for almost all frequencies (Fig. 8).

The problem with Eq. 13 is that it is based on s-wave energy gap at $T=0$ K. These two assumptions imply that the scattering rate must be zero below $2\Delta$, which is never the case with cuprates because of the d-wave gap and because the data was taken at finite temperature. In spite of this, Eq. 13 is useful because it can provide some insight into charge dynamics in cuprates. Fig. 9 displays calculations of scattering rate based on model.
spectral function $\alpha^2 F(\omega)$ shown with thin line. When the gap is zero the data qualitatively looks like underdoped YBa$_2$Cu$_3$O$_{6.95}$: at higher frequencies it is linear and it is suppressed below certain energy (black line). However for the finite values of the gap ($\Delta=200\,\text{cm}^{-1}$) the data looks more like optimal YBa$_2$Cu$_3$O$_{6.95}$: there is overshoot just above the suppressed region (gray line).

Based on these model calculations it appears that the response of YBCO on the underdoped side is dominated by coupling to bosonic mode, whereas at optimal doping the gap plays more prominent role. Indeed recent ARPES and tunneling measurements have shown that the Fermi surface of cuprates is continuously destroyed with underdoping. $^{36,37}$ On the underdoped side antinodal states do not exist (they are incoherent) and the IR response is dominated by nodal states which are coherent and not gaped. On the other hand the IR response at optimal doping is more complicated, because both antinodal (gaped) and nodal (not gaped) states are coherent and contribute to the IR response.

VII. ELECTRON–BOSON SPECTRAL FUNCTION OF Bi2212

In this section we analyze the temperature dependence of the spectral function for optimally doped Bi2212 with $T_c=91\,\text{K}$. The same data set has been analyzed before, using Eq. (3). The calculated spectra (Fig. 10) look qualitatively similar to those obtained on Y123 (Fig. 2), with a strong peak in the far-IR range followed by a dip, and high frequency contribution that extends up to several thousand cm$^{-1}$. To achieve similar levels of smoothing at different temperatures “horizontal cut” has been made (Fig. 4A).

In the normal state at $T=100\,\text{K}$ we identify a peak at $\approx 400\,\text{cm}^{-1}$ (50 meV). Note that the peak is at somewhat lower energy then in Ref. $24$, which can be traced back to the use of Eq. (3), which is strictly speaking valid only at $T=0\,\text{K}$. We also note that the peak is observed above $T_c$, unlike the $(\pi,\pi)$ resonance detected in INS only in the superconducting state. $^{46}$

As temperature decreases below $T_c$ the peak shifts to higher energies: $430\,\text{cm}^{-1}$ at $80\,\text{K}$, $520\,\text{cm}^{-1}$ at $50\,\text{K}$ and $560\,\text{cm}^{-1}$ at $10\,\text{K}$. At the lowest temperature the spectral function is almost identical to previously reported $^{26}$, which confirms that at $10\,\text{K}$ Eqs. (3) and (4) are equivalent. According to theoretical considerations in the superconducting state the peak should be off-set from the resonance frequency of the $(\pi,\pi)$ peak ($\omega_s \approx 43\,\text{meV}$) by one or two gap values ($\Delta=34\,\text{meV}$, Ref. $28$). At $10\,\text{K}$ the peak is at $70\,\text{meV}$, somewhat lower than $\Delta + \omega_s = 77\,\text{meV}$.
VIII. INVERSION OF ARPES DATA

Recently it has been argued based on ARPES data\textsuperscript{12,13,14,15} that in cuprates electrons are strongly coupled to phonons and that such strong coupling might be responsible for high $T_c$. In light of these suggestions there have been several attempts to determine $\alpha^2 F(\omega)$ from ARPES data\textsuperscript{18,38,39}. Inversion by Verga et al\textsuperscript{38} was based on the imaginary part of the self-energy $\Sigma_2(\omega)$, obtained from the real part $\Sigma_1(\omega)$ through Kramers–Kronig transformation. The spectral function was then calculated by differentiation of $\Sigma_2(\omega)$, a procedure which necessarily requires smoothing “by hand”. On the other hand Schachinger et al\textsuperscript{39} have modeled the spectral function with analytical functions and then used these models to simultaneously fit both the IR and ARPES spectra. Maximum Entropy Method (MEM) has recently been used to invert ARPES data and obtain $\alpha^2 F(\omega)$ for beryllium surface Be(10\,10) (Ref.\textsuperscript{13}) and LSCO (Ref.\textsuperscript{14}). Here we apply the same inversion method we used for IR to ARPES. The procedure of extracting $\alpha^2 F(\omega)$ is based on standard expression for the real part of quasiparticle self-energy $\Sigma_1(\omega)$ (Ref.\textsuperscript{11}):

$$\Sigma_1(\omega) = \int_0^\infty d\Omega \alpha^2 F(\Omega) \cdot \Re \left[ \Psi \left( \frac{1}{2} + i \frac{\Omega - \omega}{2\pi T} \right) - \Psi \left( \frac{1}{2} - i \frac{\Omega + \omega}{2\pi T} \right) \right],$$

where $\Psi(x)$ is digamma function. The real part of the self-energy $\Sigma_1(E_k)$ can be obtained from ARPES data as\textsuperscript{14}:

$$\Sigma_1(E_k) = E_k - \epsilon_k,$$

where $E_k$ is the renormalized dispersion measured in ARPES experiments and $\epsilon_k$ is the bare electron dispersion. As the latter function is not independently known, a common procedure when using Eq.\textsuperscript{14} is to assume a linear bare dispersion ($\epsilon_k \sim k$) and no renormalization at higher energies, i.e. $E_k = \epsilon_k$ above $\approx 250$ meV. Expression\textsuperscript{14} is again a Fredholm integral equation of the first kind and the same numerical technique described in Section\textsuperscript{11} can be used for its solution. Similar to IR, “by hand” smoothing of the data is not needed, as SVD procedure will allow us to smooth the solution by reducing the number of non-zero s.v. Since the resolution of ARPES data is poorer than IR, in all calculations we used vectors and matrices with dimensions 100 instead of 300.

As an example of this procedure in Figure\textsuperscript{11} we first present spectral function $\alpha^2 F(\omega)$ calculated from ARPES data for molybdenum surface Mo(110) (Ref.\textsuperscript{12}). As before, left panels show the calculated spectral function and right panels measured ARPES dispersion $E_k$ and dispersion calculated from Eq.\textsuperscript{14} $E_k,cal$ using the corresponding spectral function on the left. The spectral function has a characteristic shape, with a strong peak at around 200 cm$^{-1}$ and weaker structure at both higher and lower frequencies. Similar to IR, position of the main peak is fairly robust against smoothing, but weaker peaks and dips are not. The dashed lines in the left-hand panels represent $\alpha^2 F(\omega)$ calculated based on band structure\textsuperscript{33}. Low data resolution and loss of information during the inversion do not allow us to resolve the fine structure in $\alpha^2 F(\omega)$ that has been predicted numerically\textsuperscript{33}. At higher energies ($\omega \gtrsim 400$ cm$^{-1}$) the spectral function is effectively zero, in accord with band structure calculations.

These relatively simple calculations for molybdenum surface Mo(110) have uncovered the limitations of inversion of ARPES data. Fine details of the spectral function, especially narrow peaks, cannot be resolved as they are convoluted in the experimental data (Eq.\textsuperscript{14}). Maximum information that can be obtained is the frequency
region where there is significant contribution to $\alpha^2 F(\omega)$. It has recently been claimed based on MEM inversion of ARPES data that the sharp peaks identified in $\alpha^2 F(\omega)$ spectra are due to specific phonon modes. Based on our calculations we speculate that it is unlikely that such fine details of the spectra could be resolved by any inversion procedure.

Figure 12 presents the data for optimally doped Bi2212 ($T_c = 91$ K) at 130 K and 70 K taken along nodal direction. Similar to IR calculations in Fig. 10 to achieve approximately the same level of smoothing different number of s.v. values were kept in calculations at different temperatures: 8 (out of 100) at 130 K and 10 at 70 K. Within the error bars the main peak does not shift with temperature: it is at 440 cm$^{-1}$ at both 130 K and 70 K. However the peak does narrow and gains strength at 70 K (Ref. 17). Below 70 K ARPES dispersion displays almost no temperature dependence. Note also that unlike IR, there seems to be less problems with negative values in ARPES $\alpha^2 F(\omega)$ calculations. In particular there is no pronounced dip following the main peak, which might be related to the fact that the APRES scans were taken along $(\pi, \pi)$ direction where the magnitude of the gap goes to zero. Another important difference compared with IR is that there is no high frequency component in ARPES: the whole contribution to $\alpha^2 F(\omega)$ is concentrated at $\omega \lesssim 750$ cm$^{-1}$.

**IX. IR–ARPES COMPARISON**

In the previous section the inversion calculations have uncovered several important differences between the spectral function extracted from IR and ARPES. In all ARPES calculations the strong dip following the main peak was absent, which we suggested was due to the absence of the gap along the $(\pi, \pi)$ symmetry direction. More importantly, there was no high frequency contribution extending up to several thousand cm$^{-1}$ in any ARPES calculations. In this section we will make an explicit comparison between IR and ARPES spectral functions and discuss their similarities and differences.

First it should be emphasized that ARPES $\alpha^2 F(\omega)$ from Eq. (14) is not the same as the IR from Eqs. (1) and (4). ARPES is a momentum resolving technique, whereas IR averages over the Brillouin zone. More importantly ARPES probes the equilibrium $\alpha^2 F(\omega)$ (single-particle property), whereas IR measures transport $\alpha^2 F(\omega)$ (two-particle property). Recently Schachinger, et al. discussed the difference and suggested that in the simplest case these two functions might differ only by a numerical factor of 2–3. Therefore it would be very instructive to directly compare the spec-
is the contribution in IR that extends up to very high energies. There is no such contribution in any ARPES data we have available (Section VIII). Therefore based on ARPES data alone one can argue that the observed contribution to $\alpha^2 F(\omega)$ is either due to phonons or spin fluctuations. On the other hand the high frequency component in always present in IR and is necessary to keep $1/\tau(\omega)$ increasing, approximately linearly with $\omega$. Figure 13 displays calculations of $\alpha^2 F(\omega)$ for YBa$_2$Cu$_3$O$_{6.6}$ with $T_c = 59$K up to almost 1 eV (Ref. 49). Both inversion with negative values (top panels) and iterative calculations with positive values (middle panels) result in spectral function with significant contributions up to $\approx 0.85$eV. If this contribution is cut off, for example at $1,000 \text{cm}^{-1}$ (bottom panels), calculated scattering rate deviates strongly from experimental data, as $1/\tau_{\text{cal}}(\omega)$ tends to saturate above $\approx 2,000 \text{cm}^{-1}$. This result argues against phonons as the origin of the structure in $\alpha^2 F(\omega)$, as phonon spectrum cannot extend up to such high frequencies. However phonon contribution below $\approx 1,000 \text{cm}^{-1}$ cannot be ruled out.

The absence of high-frequency contribution in ARPES is puzzling and seems to indicate that the difference between IR and ARPES spectral function might be more than just a numerical prefactor. On the other hand it may also signal intrinsic problems with our procedure of extracting $\Sigma(\epsilon_k)$ from ARPES dispersion. As mentioned in Section VIII bare electron dispersion $\epsilon_k$ is not known and some assumptions must be made before Eq. (15) can be used. The most common assumptions are: 1) linear bare dispersion $\epsilon_k$ and 2) no renormalization above certain cut-off frequency. We have employed these assumptions in all our calculations, with a cut-off of typically $\approx 250 \text{meV}$. The use of both of these assumptions in highly unconventional systems like cuprates is questionable and requires further theoretical treatment.

In order to check the effect upper cut-off energy has on the solution we have performed $\alpha^2 F(\omega)$ inversion for optimally doped Bi2212 (Fig. 13) assuming that the renormalization persist up to 0.5 eV, instead of 0.25 eV. Fig. 13 also shows this new calculation with dashed line and obviously there is very little difference: the main peak is in good agreement and there is no significant contribution above $\approx 800 \text{cm}^{-1}$, even though the renormalization extends up to 0.5 eV. We speculate that in order to obtain spectral function similar to IR, either the renormalization must persist up to several eV or some more sophisticated form of the bare dispersion $\epsilon_k$ must be used.

X. SUMMARY AND OUTLOOK

A new numerical procedure of extracting electron–boson spectral function from IR and ARPES data based on inverse theory has been presented. The new method eliminates the need for differentiation and smoothing “by hand”. However we also showed that the information
is convoluted and fine details of $\alpha^2 F(\omega)$ cannot be extracted, no matter what numerical technique one uses. This especially holds for ARPES, whose current data resolution is particularly poor compared to IR.

Using this new procedure we have extracted $\alpha^2 F(\omega)$ from IR and/or ARPES data in a series of Y123 and Bi2212 samples. The calculations have uncovered several important differences between IR and ARPES spectral functions. All IR spectral functions contain, in addition to a strong peak at low frequencies ($\omega \lesssim 500$ cm$^{-1}$), contributions that extend up to very high energies (typically several thousand cm$^{-1}$). On the other hand none of ARPES spectral functions display such high energy contribution. Therefore we concluded that based on ARPES results one cannot distinguish between phonon and magnetic scenarios, as the main peak in $\alpha^2 F(\omega)$ can have either (or both) magnetic or phonon components. However in all IR results the observed high frequency contribution extends to much higher than typical phonon frequencies, the result which argues against phonon mechanism.

Finally, the observed differences between IR and ARPES have prompted us to speculate that $\alpha^2 F(\omega)$ from these two experimental techniques might contain qualitatively different information. Alternatively, we suggest that the whole concept of coupling of charge carriers to collective boson modes in the cuprates needs to be revised.

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