Optical Conductivity in a Two – Dimensional Model of the Pseudogap State

M.V.Sadovskii, N.A.Strigina
Institute for Electrophysics, Russian Academy of Sciences, Ural Branch, Ekaterinburg, 620016, Russia
E-mail: sadovski@iep.uran.ru, strigina@iep.uran.ru

Abstract

We consider a two – dimensional model of the pseudogap state, based on the scenario of strong electron scattering by fluctuations of “dielectric” (AFM, CDW) short – range order. We construct a system of recurrence equations both for one – particle Green’s function and vertex part, describing electron interaction with an external field, which take into account all Feynman graphs for electron scattering by short – range order fluctuations. The results of detailed calculations of optical conductivity are presented for different geometries (topologies) of the Fermi surface, demonstrating both the effects of pseudogap formation and localization effects. These results are in qualitative agreement with experimental data obtained for high – temperature superconducting cuprates.

PACS numbers: 74.25-q, 74.25.Jb, 74.72-h, 74.20.Mn
I. INTRODUCTION

One of the main problems in the physics of high – temperature copper oxide superconductors (HTSC) remains the nature of the so called pseudogap state [1,2], existing in the wide region of the phase diagram. Pseudogap anomalies in electronic properties of these systems are observed in a number of experiments, such as measurements of optical conductivity, NMR, inelastic neutron scattering and angle – resolved photoemission (ARPES) [1,2].

In our opinion [2], the preferable scenario for the pseudogap formation in copper oxides is based on the picture of strong scattering of current carriers on well developed fluctuations of “dielectric” (antiferromagnetic (AFM) or charge density wave (CDW)) short – range order, existing in the appropriate region of the phase diagram. This scattering is strong in the vicinity of a characteristic scattering vector \( Q = (\pi, \pi) \) (\( a \) – is the lattice spacing of the two – dimensional lattice), corresponding to the doubling of lattice period (vector of antiferromagnetism) and is a “precursor” of the spectrum transformation due to the appearance of the long – range AFM – order. Accordingly, in this region of the phase diagram develops essentially non – Fermi liquid like renormalization of electronic spectrum in certain parts of the momentum space, close to the “hot spots” on the Fermi surface [4], where this surface is effectively “destroyed”. Direct experimental confirmation of this scenario for the pseudogap formation was obtained in recent ARPES – experiments on \( Nd_{1.85}Ce_{0.15}CuO_4 \) [3], where the pseudogap renormalization of electronic spectrum was observed in the vicinity of separate “hot spots”.

Within this scenario of the pseudogap formation we can formulate a simplified “nearly exactly” solvable model, describing the main anomalies of this state [2], which takes into account all Feynman diagrams of perturbation theory for scattering by (Gaussian) fluctuations of short – range order with characteristic scattering vector from the vicinity of \( Q \), determined by the appropriate correlation length \( \xi \) [4,5]. This model is based on the two – dimensional generalization of the pseudogap formation in one – dimension due to fluctuations of CDW – type short – range order proposed rather long ago by one of the authors [6,7]. Simplified variant of this two – dimensional model (“hot patches” model) was used in Refs. [8–11] to describe basic anomalies of superconducting state, formed on the “background” of the “dielectric” pseudogap.

In Refs. [4,5] mainly one – particle properties of the model were analyzed, such as the spectral density and the density of states. The remarkable property of this model is the possibility to sum also the whole Feynman series for the two – particle problem for the vertex part, describing the system response to an external field (e.g. electromagnetic) [6,12,13]. In the simplified version of the “hot patches” model such calculations for the two – dimensional case were performed in Ref. [14]. The main aim of the present work is the detailed analysis of both the theoretical aspects of calculations of two – particle properties of the more general model [4,5] and calculations of the optical conductivity for different geometries (topologies) of the Fermi surface, appearing for “realistic” enough spectrum of non – interacting electrons.

II. “HOT SPOTS” MODEL.
1. Description of the model and “nearly exact” solution for the one–particle Green’s function.

In the model of “nearly antiferromagnetic” Fermi–liquid, which is actively used to explain the microscopic nature of HTSC \[15,16\], effective electron interaction with spin–fluctuations of AFM short–range order is usually described by the dynamic spin susceptibility \(\chi_\mathbf{q}(\omega)\) of the following form, determined by fitting to NMR data \[16\]:

\[
V_{\text{eff}}(\mathbf{q}, \omega) = g^2 \chi_\mathbf{q}(\omega) \approx g^2 \frac{\xi^2}{1 + \xi^2 (\mathbf{q} - \mathbf{Q})^2 - \frac{i \omega}{\omega_{sf}}} \tag{1}
\]

where \(g\) – is a coupling constant, \(\xi\) – correlation length of spin fluctuations, \(\mathbf{Q} = (\pi/a, \pi/a)\) – vector of antiferromagnetic ordering in dielectric phase, \(\omega_{sf}\) – characteristic frequency of spin fluctuations. Both dynamic susceptibility and effective interaction (1) have a sharp maximum in the region of \(\mathbf{q} \sim \mathbf{Q}\), leading to two “types” of quasiparticles — “hot” one with momenta in the vicinity of “hot spots” on the Fermi surface (Fig. 1) and “cold” with momenta around the parts of the Fermi surface around Brillouin zone diagonals \[4\]. This is due to the fact that quasiparticles from the regions around the “hot spots” are strongly scattered with the momentum transfer of the order of \(\mathbf{Q}\) by spin fluctuations (1), while for quasiparticles with momenta far from the “hot spots” this interaction is small enough.

For high enough temperatures \(\pi T \gg \omega_{sf}\) we can neglect the spin dynamics \[4\], and use the static approximation in (1):

\[
V_{\text{eff}}(\mathbf{q}) = \tilde{\Delta}^2 \frac{\xi^2}{1 + \xi^2 (\mathbf{q} - \mathbf{Q})^2} \tag{2}
\]

where \(\tilde{\Delta}\) – is an effective parameter with dimension of energy. In the following, similar to Refs. \[4,5\], we consider both \(\Delta\) and \(\xi\) as phenomenological parameters, to be determined from experiments. Considerable simplification of calculations, allowing to analyze higher order contributions of perturbation theory, can be achieved if we replace (2) by the model interaction of the following form \[5\]:

\[
V_{\text{eff}}(\mathbf{q}) = \Delta^2 \frac{2 \xi^{-1}}{\xi^{-2} + (q_x - Q_x)^2 \xi^{-2} + (q_y - Q_y)^2} \tag{3}
\]

where \(\Delta^2 = \tilde{\Delta}^2/4\). Eq. (3) is qualitatively the same as (2) and quantitatively is very close to it in most interesting region of \(|\mathbf{q} - \mathbf{Q}| < \xi^{-1}\), determining the scattering in the vicinity of “hot spots”.

We assume the following spectrum of free (“bare”) quasiparticles \[4\]:

\[
\xi_p = -2t(\cos p_x a + \cos p_y a) - 4t' \cos p_x a \cos p_y a - \mu \tag{4}
\]

where \(t\) – is the transfer integral between nearest neighbors, while \(t'\) – is the transfer integral for second nearest neighbors on the square lattice, \(\mu\) – is the chemical potential. This expression gives rather good approximation to the results of band structure calculations for real HTSC – systems, e.g. for \(YBa_2Cu_3O_{6+\delta}\) we have \(t = 0.25eV\), \(t' = -0.45t\) \([4]\). Chemical potential \(\mu\) is fixed by carrier concentration. In this paper we shall consider different characteristic values of \(t\), \(t'\) and \(\mu\), leading to different geometries (topologies) of
Fermi surfaces, with no relation to any particular system, with the aim of analyzing general enough picture.

Consider first – order in $V_{\text{eff}}$ correction to electronic self – energy, determined by the simplest Feynman diagram ($\varepsilon_n = (2n + 1)\pi T$):

$$\Sigma(\varepsilon_n\mathbf{p}) = \sum_{\mathbf{q}} V_{\text{eff}}(\mathbf{q}) \frac{1}{\varepsilon_n - \xi_{\mathbf{p}+\mathbf{q}}} \quad (5)$$

For large enough correlation lengths $\xi$, the main contribution to the sum over $\mathbf{q}$ comes from the region close to $\mathbf{Q} = (\pi/a, \pi/a)$. Then we can write:

$$\xi_{\mathbf{p}+\mathbf{q}} = \xi_{\mathbf{p}+\mathbf{Q}+\mathbf{k}} \approx \mathbf{v}_{\mathbf{p}+\mathbf{Q}} \mathbf{k} \quad (6)$$

where $\mathbf{v}_{\mathbf{p}+\mathbf{Q}} = \frac{\partial \xi_{\mathbf{p}+\mathbf{Q}}}{\partial \mathbf{p}}$ – is the appropriate velocity of the quasiparticle on the Fermi surface. Then (5) is easily calculated to be:

$$\Sigma(\varepsilon_n\mathbf{p}) = \frac{\Delta^2}{\varepsilon_n - \xi_{\mathbf{p}+\mathbf{Q}} + i(\varepsilon_{\mathbf{p}+\mathbf{Q}}^x + \varepsilon_{\mathbf{p}+\mathbf{Q}}^y)\kappa \text{sign} \varepsilon_n} \quad (7)$$

where $\kappa = \xi^{-1}$. Let us stress that “linearization” of quasiparticle spectrum (6) in (5) is done only for a small (of the order of $v_F \xi^{-1}$), due to a large values of $\xi$, correction to an energy spectrum of quasiparticles close to the Fermi surface, while the form of the spectra $\xi_{\mathbf{p}}$ and $\xi_{\mathbf{p}+\mathbf{Q}}$ is given by general expression (4).

In Ref. [5] we have performed a detailed analysis of higher – order contributions for $\Sigma(\varepsilon_n\mathbf{p})$. It was shown that in case of equal signs of velocity projections (on the Fermi surface) $v_{\mathbf{p}}^x$ and $v_{\mathbf{p}+\mathbf{Q}}^x$, as well as of $v_{\mathbf{p}}^y$ and $v_{\mathbf{p}+\mathbf{Q}}^y$, Feynman integrals for any diagram of any order are determined only by contributions from the poles of Lorentzians in (3) and can be easily calculated [1]. In this case the contribution of an arbitrary diagram of the $N$-th order over effective interaction (3) is given by:

$$\Sigma^{(N)}(\varepsilon_n\mathbf{p}) = \Delta^{2N} \prod_{j=1}^{2N-1} \frac{1}{\varepsilon_n - \xi_j(\mathbf{p}) + in_j v_j \kappa} \quad (8)$$

where $\xi_j(\mathbf{p}) = \xi_{\mathbf{p}+\mathbf{Q}}$ and $v_j = |v_{\mathbf{p}+\mathbf{Q}}^x| + |v_{\mathbf{p}+\mathbf{Q}}^y|$ for odd $j$, while $\xi_j(\mathbf{p}) = \xi_{\mathbf{p}}$ and $v_j = |v_{\mathbf{p}}^x| + |v_{\mathbf{p}}^y|$ for even $j$. Here $n_j$ – is the number interaction lines surrounding $j$-th Green’s function in a given diagram and for definiteness we assumed $\varepsilon_n > 0$.

In Ref. [5] we have given a detailed analysis of conditions for these requirements for velocities in the points on the Fermi surface, connected by vector $\mathbf{Q}$ (“hot spots”) to be valid, and also presented some examples of appropriate geometries of the Fermi surface, which are realized for certain relations between transfer integrals $t$ and $t'$ in Eq. (4). Under these conditions expression (8) is exact, with only limitation due to the use of abovementioned “linearization” in Eq. (6). In general case (for other relations between $t$ and $t'$) we use

---

1 Analogous statement is valid also for a special case of velocities in “hot spots”, connected by vector $\mathbf{Q}$, being exactly perpendicular to each other [4].
as a fortunate Ansatz for the contribution of an arbitrary order, obtained by simple continuation over spectrum parameters \( t \) and \( t' \) to the region of interest. Even in the worst one-dimensional case [7], equivalent to the case of the simple square Fermi surface, appearing in two dimensions for \( t' = 0 \) and \( \mu = 0 \), the use of this Ansatz gives results (e.g. for the density of states) which are quantitatively very close [17] to the results of an exact numerical simulation of this problem [18]. In this sense we are using the term “nearly exact” solution.

In case of the use of this Ansatz Eq. (8) it is easily seen, that a contribution of some arbitrary diagram with crossing interaction lines is in fact equal to the contribution of some definite diagram of the same order without intersections of interaction lines [7]. Thus we can limit ourselves only to diagrams without crossing interaction lines, taking those with intersections into account with the help of additional combinatorial factors, attributed to “initial” interaction vertices (or interaction lines) [7]. As a result we obtain the following recursion relation for the one-electron Green’s function (continuous fraction representation) [7], giving an effective algorithm for numerical computations [3]:

\[
G_k(\varepsilon_n \xi_p) = \frac{1}{i\varepsilon_n - \xi_k(p) + ikv_k\kappa - \Sigma_{k+1}(\varepsilon_n \xi_p)} \\
\equiv \left\{ G_{0k}(\varepsilon_n \xi_p) - \Sigma_{k+1}(\varepsilon_n \xi_p) \right\}^{-1}
\]  

\[\Sigma_k(\varepsilon_n \xi_p) = \Delta^2 \frac{v(k)}{i\varepsilon_n - \xi_k(p) + ikv_k\kappa - \Sigma_{k+1}(\varepsilon_n \xi_p)} \]  

In graphic form this recursion relation for the Green’s function is shown in Fig. 2. The physical Green’s function of interest to us is given by \( G(\varepsilon_n \xi_p) = G_{k=0}(\varepsilon_n \xi_p) \). In (9) we also introduced a helpful technical notation:

\[
G_{0k}(\varepsilon_n \xi_p) = \frac{1}{i\varepsilon_n - \xi_k(p) + ikv_k\kappa}
\]

Combinatorial factor:

\[
v(k) = k
\]

for our case of commensurate fluctuations with \( Q = (\pi/a, \pi/a) \) [4], if we neglect the spin variables (CDW – fluctuations). If we take into account the spin structure of interaction within the model of “nearly antiferromagnetic” Fermi liquid (spin – fermion model of Ref. [4]), combinatorics of diagrams becomes more complicated. In this case, spin – conserving scattering formally gives commensurate combinatorics, while spin – flip scattering is described by diagrams with incommensurate combinatorics (“charged” random field in terms of Ref. [4]). As a result, the recursion relation for the Green’s function (10) is still valid, but with different combinatorial factor \( v(k) \) [4]:

\[
v(k) = \begin{cases} 
\frac{k+2}{3} & \text{for odd } k \\
\frac{k}{3} & \text{for even } k
\end{cases}
\]

Below we limit ourselves to the cases of (12) and (13), some details concerning the case of incommensurate CDW – fluctuations can be found in Ref. [3 7].
Our solution for one-particle Green’s function is exact in the limit of $\xi \to \infty$, when it can be found also in analytic form \cite{1,4}. It is also exact in a trivial limit of $\xi \to 0$, when for fixed value of $\Delta$ interaction \cite{4} just vanishes. For all intermediate values of $\xi$ it gives apparently very good interpolation, being practically exact for certain geometries of the Fermi surface, which are realized for definite regions of spectrum parameters in \cite{1} \cite{5}.

Using (14) we can easily perform numerical computations of one-electron spectral density:

$$A(Ep) = -\frac{1}{\pi} ImG^R(Ep)$$

which can be determined also from ARPES – experiments \cite{2}. In (14) $G^R(Ep)$ denotes the retarded Green’s function, which is obtained by the usual analytic continuation of (9) from Matsubara’s frequencies to the real axis of $E$. Similarly, we can perform calculations of one-electron density of states:

$$N(E) = \sum_p A(Ep) = -\frac{1}{\pi} \sum_p ImG^R(Ep)$$

Details of these calculations and discussion of results for our two-dimensional model can be found in Refs. \cite{4,5}.

2. Recurrence relations for the vertex part and conductivity.

To calculate optical conductivity we need the knowledge of the vertex part describing electromagnetic response of the system. This vertex can be determined using the method suggested for the similar one-dimensional problem in Refs. \cite{12,13}. Arbitrary diagram for the vertex part can be obtained by an insertion of an external field line to the appropriate diagram for the self-energy \cite{6}. We have already noted that in our model we can limit ourselves only to diagrams with non-intersecting interaction lines with additional combinatorial factors $v(k)$ in “initial” interaction vertices. It is clear then that to calculate vertex corrections we have to consider only diagrams of the type shown in Fig. 4. Then we immediately obtain the system of recurrence equations for the vertex parts shown by diagrams of Fig. 4. To find appropriate analytic expressions consider the simplest vertex correction shown in Fig. 3(a). Performing explicit calculations for $T = 0$ in $RA$ – channel we find its contribution to be:

$$J^{(1)RA}_1(\varepsilon; \varepsilon + \omega; p + q) = \sum_K V_{eff}(K)G^A_{00}(\varepsilon; \xi_p - K)G^R_{00}(\varepsilon + \omega; \xi_{p+q}) =$$

$$= \Delta^2 \left\{ G^A_{00}(\varepsilon; \xi_1(p) + iv_1K) - G^R_{00}(\varepsilon + \omega; \xi_1(p + q) - iv_1K) \right\} \frac{1}{\omega + \xi_1(p) - \xi_1(p + q)} =$$

$$= \Delta^2 G^A_{00}(\varepsilon; \xi_1(p) + iv_1K)G^R_{00}(\varepsilon + \omega; \xi_1(p + q) - iv_1K) \left\{ 1 + \frac{2iv_1K}{\omega + \xi_1(p) - \xi_1(p + q)} \right\} \equiv \Delta^2 G^A_{01}(\varepsilon; \xi_p)G^R_{01}(\varepsilon + \omega; \xi_{p+q}) \left\{ 1 + \frac{2iv_1K}{\omega + \xi_1(p) - \xi_1(p + q)} \right\}$$

(16)
where during the integral calculations we have used the following identity, valid for the free electron Green’s functions:

\[ G^A_{00}(\varepsilon \xi_p)G^R_{00}(\varepsilon + \omega \xi_{p+q}) = \left\{ G^A_{00}(\varepsilon \xi_p) - G^R_{00}(\varepsilon + \omega \xi_{p+q}) \right\} \frac{1}{\omega - \xi_{p+q} + \xi_p} \]  \hspace{1cm} (17)

“Dressing” the internal electronic lines we obtain the diagram shown in Fig. 2(b), so that using the identity:

\[ G^A(\varepsilon \xi_p)G^R(\varepsilon + \omega \xi_{p+q}) = \left\{ G^A(\varepsilon \xi_p) - G^R(\varepsilon + \omega \xi_{p+q}) \right\} \times \frac{1}{\omega - \xi_{p+q} + \xi_p - \Sigma_i^R(\varepsilon + \omega \xi_{p+q}) + \Sigma_i^A(\varepsilon \xi_p)} \]  \hspace{1cm} (18)

valid for exact Green’s functions, we can write the contribution of this diagram as:

\[ J^{RA}_1(\varepsilon \xi_p; \varepsilon + \omega \xi_{p+q}) = \Delta^2 v(1)G^A_1(\varepsilon \xi_p)G^R_1(\varepsilon + \omega \xi_{p+q}) \times \left\{ 1 + \frac{2i v_i \kappa}{\omega - \xi_1(\varepsilon \xi_p) - \xi_1(\varepsilon \xi_p) - \Sigma_2^R(\varepsilon + \omega \xi_{p+q}) + \Sigma_2^A(\varepsilon \xi_p)} \right\} J^{RA}_1(\varepsilon \xi_p; \varepsilon + \omega \xi_{p+q}) \]  \hspace{1cm} (19)

Here we have assumed that interaction line in the vertex correction of Fig. 2(b) “transforms” self – energies \( \Sigma_i^{R,A} \) of internal lines into \( \Sigma_i^{R,A} \), in accordance with our main approximation for the self – energy (cf. Fig. 2)

Now we can write down the similar expression for the general diagram, shown in Fig. 3(c):

\[ J^{RA}_k(\varepsilon \xi_p; \varepsilon + \omega \xi_{p+q}) = \Delta^2 v(1)G^A_k(\varepsilon \xi_p)G^R_k(\varepsilon + \omega \xi_{p+q}) \times \left\{ 1 + \frac{2i v_k \kappa}{\omega - \xi_k(\varepsilon \xi_p) - \xi_k(\varepsilon \xi_p) - \Sigma_{k+1}^R(\varepsilon + \omega \xi_{p+q}) + \Sigma_{k+1}^A(\varepsilon \xi_p)} \right\} J^{RA}_k(\varepsilon \xi_p; \varepsilon + \omega \xi_{p+q}) \]  \hspace{1cm} (20)

Then we can write the general recurrence relation for the vertex part Fig. 2 in the following form:

\[ J^{RA}_{k-1}(\varepsilon \xi_p; \varepsilon + \omega \xi_{p+q}) = 1 + \Delta^2 v(1)G^A_k(\varepsilon \xi_p)G^R_k(\varepsilon + \omega \xi_{p+q}) \times \left\{ 1 + \frac{2i v_k \kappa}{\omega - \xi_k(\varepsilon \xi_p) - \xi_k(\varepsilon \xi_p) - \Sigma_{k+1}^R(\varepsilon + \omega \xi_{p+q}) + \Sigma_{k+1}^A(\varepsilon \xi_p)} \right\} J^{RA}_k(\varepsilon \xi_p; \varepsilon + \omega \xi_{p+q}) \]  \hspace{1cm} (21)

The “physical” vertex \( J^{RA}(\varepsilon \xi_p; \varepsilon + \omega \xi_{p+q}) \) is determined as \( J^{RA}_{k=0}(\varepsilon \xi_p; \varepsilon + \omega \xi_{p+q}) \). Recurrence procedure (21) takes into account all perturbation theory diagrams for the vertex part. For \( \kappa \to 0 \) \( (\xi \to \infty) \) (21) reduces to the series studied in Ref. 8 (cf. also Ref. 11), which can be summed exactly in analytic form. Standard “ladder” approximation corresponds in our scheme to the case of combinatorial factors \( v(k) \) in (21) being equal to 1 13.

Conductivity of our system can be expressed via retarded density – density response function \( \chi^R(q\omega) \):

\[ \chi^R(q\omega) \]

---

2One of the main motivations for this trick is that it guarantees the fulfillment of an exact Ward identity to be discussed below.
\[
\sigma(\omega) = e^2 \lim_{q \to 0} \left( -\frac{i\omega}{q^2} \right) \chi^R(q\omega) \tag{22}
\]
where \(e\) is electronic charge,

\[
\chi^R(q\omega) = \omega \left\{ \Phi^{RA}(0q\omega) - \Phi^{RA}(00\omega) \right\} \tag{23}
\]
while two-particle Green’s function \(\Phi^{RA}(\varepsilon q\omega)\) is determined by the loop graph shown in Fig. [x].

Direct numerical computations confirm that the recurrence procedure \((21)\) satisfies an exact relation, which directly follows (for \(\omega \to 0\)) from the Ward identity derived in Ref. \([19]\):

\[
\Phi^{RA}(00\omega) = -\frac{N(E_F)}{\omega} \tag{24}
\]
where \(N(E_F)\) is the density of states at the Fermi level \(E_F = \mu\). This is the main argument for the validity of an Ansatz used to derive Eqs. \((19), (20)\) and \((21)\).

Finally, we can write conductivity in the following symmetrized form, convenient for numerical computations:

\[
\sigma(\omega) = \frac{e^2\omega^2}{\pi} \lim_{q \to 0} \frac{1}{q^2} \sum_p \left\{ G^R \left( \frac{\omega}{2}, p + \frac{q}{2} \right) J^{RA} \left( \frac{\omega}{2}, p + \frac{q}{2}; -\frac{\omega}{2}, p - \frac{q}{2} \right) G^A \left( -\frac{\omega}{2}, p - \frac{q}{2} \right) - 
- G^R \left( \frac{\omega}{2}, p \right) J^{RA} \left( \frac{\omega}{2}, p; -\frac{\omega}{2}, p \right) G^A \left( -\frac{\omega}{2}, p \right) \right\} \tag{25}
\]
where we have introduced an additional factor of 2 due to spin summation.

Direct numerical computations were performed using \((25), (21), (9)\), with recurrence procedure being cut for high enough \(k\), where all \(\Sigma_k\) are \(J_k\) were supposed to be equal to zero. Integration in \((25)\) was made over the whole Brillouin zone. The “bare” electronic spectrum was taken from \((4)\). Integration momenta are made dimensionless in a natural way with the help of the lattice constant \(a\), while all energies in what follows are in units of the transfer integral \(t\). Conductivity is measured in units of the universal conductivity in two-dimensions \(\sigma_0 = \frac{e^2}{\hbar} = 2.5 \times 10^{-4} \text{ Ohm}^{-1}\), and the density of states — in units of \(1/ta^2\).

III. RESULTS AND DISCUSSION.

Optical conductivity and some other characteristics of the model were computed for different values of parameters, determining the “bare” quasiparticle spectrum \((4)\) and for fixed value of \(\Delta = t\). Let us consider first Fermi surfaces close to the case of half-filled band with \(\mu = 0\) and \(t' = 0\), shown (for the first quadrant of the Brillouin zone) in Fig. [7] (a). For \(\mu = 0\) and \(t' = 0\) the Fermi surface is a well known simple square (complete nesting), so that this case is quite similar to that of one-dimensional system, considered in Refs. \([8,12,13]\). Our results for the real part of optical conductivity of two-dimensional model, for the case of spin-fermion combinatorics of diagrams and different values of
correlation length of AFM short – range order (parameter \( \kappa = \xi^{-1} \), where \( \xi \) is measured in units of lattice spacing \( a \)) are shown in Fig. 8. Qualitative form of conductivity is similar to that found previously for the one – dimensional model (for the case of incommensurate CDW – fluctuations) in Refs. [12,13]. It is characterized by a sharp maximum due to pseudogap absorption (corresponding densities of states, demonstrating pseudogap at the Fermi level, are shown at the insert in Fig. 8) at \( \omega \sim 2\Delta \) and also by weaker maximum at smaller frequencies due to Anderson localization of carriers in the random field of static AFM fluctuations. Localization nature of this maximum is confirmed by its transformation into the usual “Drude – like” peak (with maximum at \( \omega = 0 \)) if we perform calculations in the “ladder” approximation, when combinatorial factors \( v(k) = 1 \), corresponding to “switching off” the contribution of diagrams with intersecting interaction lines, leading to two – dimensional Anderson localization [19,20]. Qualitative form of conductivity in this case is also quite similar to that obtained previously for one – dimensional model in Ref. [13].

The narrowing of localization peak with diminishing correlation length of fluctuations can be explained, as was noted in Ref. [13], by weakening of effective interaction (3) for smaller values of \( \xi \) (for the fixed value of \( \Delta \)), leading to general weakening of scattering rate also at the “cold” parts of the Fermi surface. Note that general behavior of our density of states and optical conductivity is in full qualitative agreement with the results of quantum Monte Carlo calculations of a similar model of two – dimensional Peierls transition obtained recently in Ref. [21].

Consider now the case of \( \mu = 0 \) again, but “switch on” the second nearest neighbor transfer integral \( t' \) in (4). Then we are dealing with Fermi surfaces different from the square one, as shown in Fig. 7 (a). At the insert in this figure we show the energy dependence of spectral densities (14) in several typical points of these Fermi surfaces. It can be seen that these spectral densities have characteristic “non – Fermi liquid” like behavior of the type studied in Refs. [4,5], practically everywhere at the Fermi surface, while this surface is not very far from the square, despite the fact that the “hot spot” in this case is precisely at the crossing of the zone diagonal with the Fermi surface. Corresponding frequency dependences of the real part of optical conductivity are shown in Fig. 9. At the insert in this figure we show corresponding densities of states. It is seen that as we depart from the situation of complete nesting, conductivity maximum due to pseudogap transitions becomes smaller, while the localization peak grows and becomes sharper (in accordance with the general sum rule for conductivity). Note, however, that pseudogap absorption peak remains noticeable even in the case, when the pseudogap in the density of states is practically absent (curves 4 in Fig. 9).

Let us now return to the case of \( t' = 0 \) and vary chemical potential \( \mu \), obtaining Fermi surfaces, which are sufficiently close to square, as shown in Fig. 7 (b). Strictly speaking there are no “hot spots” at these Fermi surfaces at all, but the spectral density, shown at the insert in Fig. 7 (b) still conserves a typical pseudogap form. Corresponding dependences for the real part of optical conductivity are shown in Fig. 10.

Consider now different geometries of the Fermi surface with “hot spots”, shown in Fig. 11. In Figs. 12, 13 we show the real part of optical conductivity, calculated (for different diagram combinatorics) for two characteristic values of \( t' = -0.4t \) and \( t' = -0.6t \) for the chemical potential value \( \mu = 0 \), when “hot spots” are on the zone diagonal (curve 5 in Fig. 11 (a) and curve 4 in Fig. 11 (b)). We can see again that the pseudogap behavior of conductivity
persists even for the case, when the pseudogap in the density of states (shown at the inserts in Figs. 12, 13) is practically absent. Dashed curve in Fig. 12 shows the results of the “ladder” approximation, demonstrating typical vanishing of two-dimensional localization in this approximation. The results shown in Fig. 13 demonstrate typical “smearing” of the pseudogap maximum of conductivity with diminishing correlation length of short-range order fluctuations.

For majority of copper oxide high-temperature superconductors typical geometry of the Fermi surface is described by the case of \( t' = -0.4t \) and \( \mu = -1.3t \) [4] (as shown by curve 3 in Fig. 11 (a)). Results of our calculations of optical conductivity for this case and for different values of the inverse correlation length \( \kappa \) are shown in Fig. 14 (for spin-fermion combinatorics of diagrams). Here we also introduced an additional weak scattering due to inelastic processes, using the standard substitution \( \omega \rightarrow \omega + i\gamma \) [22], which leads to the appearance of a narrow “Drude-like” peak in the region of \( \omega < \gamma \) (destruction of two-dimensional localization due to dephasing). It can be easily checked that for higher values of inelastic scattering rate \( \gamma \) localization peak is “smeared” and transforms into the “usual” Drude-like peak in the region of small frequencies. Pseudogap absorption maximum becomes more pronounced with the growth of correlation length \( \xi \) (diminishing \( \kappa \)). In Fig. 15 we show frequency dependencies of an effective scattering rate \( 1/\tau(\omega) \) and effective mass \( m^*(\omega) \), which can be determined from our calculations, using the generalized Drude formula, which is often used to fit experimental data [1]:

\[
\frac{1}{\tau(\omega)} = \frac{\omega_p^2}{4\pi} Re \left( \frac{1}{\sigma(\omega)} \right)
\]

(26)

\[
\frac{m^*(\omega)}{m} = -\frac{1}{2\omega_p^2} Re \left( \frac{1}{\sigma(\omega)} \right)
\]

(27)

Here \( \omega_p \) – is plasma frequency, \( m \) – is the mass of a free electron. From Fig. 13 we can see, that \( 1/\tau(\omega) \) (expressed in this figure in units of \( \frac{\omega_p^2}{4\pi e^2 h} \)) demonstrates quite typical pseudogap behavior for the frequency range of \( \omega < 2\Delta \). Note, that the density of states in this case possess only rather weak pseudogap [5] (cf. insert in Fig. 13). In Fig. 16 we show similar results for the same case (typical for HTSC – oxides), but for commensurate combinatorics of diagrams (CDW – type fluctuations). We see that in this case the pseudogap absorption maximum is almost invisible.

From Fig. 14 (b) we can see that with the change of the chemical potential in the interval from \( \mu = 0 \) to \( \mu = -1.666t \) the Fermi surface acquires larger and larger “flat” parts, transforming for \( \mu \approx 1.666t \) practically into a “cross-like”. Similar Fermi surface was observed in ARPES experiments on \( La_{1.28}Nd_{0.6}Sr_{0.12}CuO_4 \) [23,24]. In this case velocity projections in “hot spots”, connected by \( Q = (\frac{\pi}{a}, \frac{\pi}{a}) \), become orthogonal. For \( \mu/t = -1.666... \) topology of the Fermi surface change (cf. Fig. 14 (b)) and in the whole region of \( \mu/t < -1.666... \) these projections have the same signs, guaranteeing the exactness of our main Ansatz [8] for the contributions of higher order diagrams [5]. It is of some interest to present the results of our calculations of optical conductivity also for these values of \( \mu \). These are shown (for the case of commensurate (CDW) diagram combinatorics) in Fig. 17, where we can follow the changes of localization peak in conductivity as the chemical potential passes through the region of topological phase transition. A weak pseudogap absorption maximum practically does not change at all. At the insert in Fig. 17 we show the evolution of localization peak
as we “switch on” inelastic scattering processes (scattering rate $\gamma$) for the case of $\mu = -1.8t$. The transition from localization to Drude–like behavior due to inelastic dephasing is clearly seen. These results show, that the change in topology of the Fermi surface does not lead to any strong qualitative change of optical conductivity in our model.

**IV. CONCLUSION**

Our analysis shows the variety of results, which may be obtained in this model for different geometries and topologies of the Fermi surface, appearing for different values of the “bare” quasiparticle energy spectrum $\varepsilon_0$. It is interesting to compare these data with results obtained earlier for the simplified model with “hot patches” on the Fermi surface [4]. Pseudogap anomalies in the “hot patches” model were determined mainly by strong scattering on these (flat) patches only, as well as by their relative size. Accordingly, the localization peak of conductivity in this model was rather weak and dominating behavior at small frequencies was determined by Drude–like peak due to the scattering on “cold” parts of the Fermi surface with the scattering rate $\gamma$ (analogous to inelastic scattering rate defined above). Analysis of the more realistic model, given in this paper, shows that the contribution of localization peak may be more pronounced and in fact this peak may transform into a narrow enough “Drude–like” peak with the inclusion of dephasing processes.

Probably the main deficiency of the present model is the neglect of the dynamics of fluctuations of short–range order. This approximation is justified, as was noted in Refs. [4], only for high enough temperatures. However, at higher temperatures inelastic scattering, responsible for dephasing processes and breakdown of localization, become more important. The other shortcoming, as was stressed several times [7,5], is our limitation to Gaussian fluctuations only of short–range order, which also may be justified only for high enough temperatures.

Discussing the possible relation of the above results with experimental data obtained on real HTSC–cuprates we note, that in majority of these experiments [2,4] localization peak was not observed at all, which may be due to the large contribution of inelastic processes (dephasing) at high enough temperatures used in these measurements. Peaks in optical conductivity at small frequencies, attributed to localization, were observed in disordered samples of $YBaCuO$ in Refs. [25,26]. In recent experiments on $NdCeCuO$ [27,28] this peak was especially clearly seen. In particular, the qualitative behavior of optical conductivity observed in Ref. [28] for several samples of $NdCeCuO$ with different compositions (from underdoped to optimally doped) is in complete agreement with our results shown above in Fig. 14, which were considered as typical for HTSC–cuprates. We conclude that the “hot spots” model may be successfully used for realistic enough description of anomalies of optical conductivity in high–temperature superconductors.

The authors are grateful to E.Z.Kuchinskii for numerous discussions. This work was supported in part by the grants of the Russian Foundation of Basic Research 02-02-16031, CRDF No. REC-005 and the program of fundamental research of the Presidium of the Russian Academy of Sciences “Quantum Macrophysics”. It was also partly supported by the project of the Russian Ministry of Industry and Science “Studies of collective and quantum effects in condensed matter”.

11
FIG. 1. Fermi surface with “hot spots”, connected by the scattering vector of the order of $Q = (\frac{\pi}{a}, \frac{\pi}{a})$. 
FIG. 2. Diagrammatic form of the recurrence relation for the Green's function.
FIG. 3. General diagram for the vertex correction.
FIG. 4. Recurrence relations for the vertex part.
FIG. 5. Simplest corrections for vertex parts.
\[ \Phi^{RA}(q \epsilon \omega) = \frac{1}{2\pi i} \sum_p \]
FIG. 7. Fermi surfaces for different values of $t'$ and chemical potential $\mu$. (a) corresponds to $\mu = 0$ and the following values of $t'/t$: 0 – I; -0.2 – II; -0.4 – III; -0.6 – IV. (b) corresponds to $t' = 0$ and following values of $\mu/t$: 0 – I; -0.3 – II; -0.5 – III; -0.6 – IV. At the inserts – energy dependences of the spectral density for spin–fermion model for $\kappa a = 0.1$ at the points in the momentum space denoted by stars.
FIG. 8. Real parts of optical conductivity in spin – fermion model for the case of square Fermi surface ($\mu = 0$, $t' = 0$) for different values of inverse correlation length of short – range order $\kappa a$: $0.1 – 1$; $0.2 – 2$; $0.5 – 3$. At the insert – appropriate densities of states.
FIG. 9. Real part of optical conductivity in spin–fermion model for $\mu = 0$ and $\kappa a = 0.1$ for different Fermi surfaces, obtained from the square by “switching on” the transfer integral $t'/t$: 0 – 1; -0.2 – 2; -0.4 – 3; -0.6 – 4. At the insert – appropriate densities of states.
FIG. 10. Real part of optical conductivity in spin–fermion model for $t' = 0$ and $\kappa a = 0.1$ for different Fermi surfaces, obtained from the square by the shift from half–filled band. Chemical potentials correspond to the following values of $\mu/t$: 0 – 1; -0.3 – 2; -0.5 – 3; -0.6 – 4.
FIG. 11. Fermi surfaces for different values of $t'$ and chemical potential $\mu$. (a) corresponds to the case of $t'/t = -0.4$, typical for HTSC–cuprates, and the following values of $\mu/t$: -1.6 – 1; -1.4 – 2; -1.3 – 3; -1.1 – 4; 0 – 5. “Hot spots” exist for $-1.6 < \mu/t < 0$. (b) corresponds to the case of $t'/t = -0.6$ and the following values of $\mu/t$: -1.8 – 1; -1.666 – 2; -1.63 – 3; 0 – 4. “Hot spots” exist for $\mu < 0$. Dashed line – magnetic Brillouin zone.
FIG. 12. Real part of optical conductivity for $t'/t = -0.4$ and $\mu/t = 0$ for $\kappa a = 0.1$ and different combinatorics of diagrams: 1 – spin – fermion combinatorics; 2 – commensurate case. Dashed line - “ladder” approximation. At the insert – corresponding densities of states.
FIG. 13. Real part of optical conductivity in spin–fermion model for the case of $t'/t = -0.6$ and $\mu = 0$ for different values of inverse correlation length $\kappa a$: $0.1 – 1$; $0.2 – 2$; $0.5 – 3$; $1 – 4$. At the insert – densities of state, corresponding to 1 and 4.
FIG. 14. Real part of optical conductivity in spin–fermion model for $t'/t = -0.4$ and $\mu/t = -1.3$ and different values of correlation length $\kappa a$: 0.05 – 1; 0.1 – 2; 0.2 – 3. Dephasing rate $\gamma/t = 0.005$. 
FIG. 15. Generalized scattering rate and effective mass for the case of $t'/t = -0.4$ and $\mu/t = -1.3$, typical for high-temperature superconductors. Parameters of the generalized Drude model were obtained for spin-fermion model with different values of correlation length $\kappa a$: 0.05 – 1; 0.1 – 2; 0.2 – 3. Dephasing rate $\gamma/t = 0.005$. 
FIG. 16. Real part of optical conductivity for commensurate case with $t'/t = -0.4$ and $\mu/t = -1.3$ and different values of inverse correlation length $\kappa a$: 0.05 – 1; 0.1 – 2; 0.2 – 3. Dephasing rate $\gamma = 0.005/t$. 
FIG. 17. Evolution of the real part of optical conductivity for commensurate case with $t'/t = -0.6$ and $\kappa a = 0.2$ and chemical potential changing in the vicinity of topological transition. Different curves correspond to the following values of $\mu/t$: -1.79 – 1; -1.77 – 2; -1.66 – 3; -1.63 – 4. Dephasing rate $\gamma/t = 0.01$. At the insert – real part of optical conductivity for the case of $\mu/t = -1.8$ for different values of $\gamma/t$: 0 – 1; 0.01 – 2; 0.05 – 3.
REFERENCES

[1] T.Timusk, B.Statt. Rep.Progr.Phys. 62, 61 (1999)
[2] M.V.Sadovskii. Usp.Fiz.Nauk 171, 539 (2001); (Phys.Uspekhi 44, 515 (2001)); ArXiv: cond-mat/0102111
[3] N.P.Armitage, D.H.Lu, C.Kim, A.Damascelli, K.M.Shen, F.Ronning, D.L.Feng, P.Bogdanov, Z.-X.Shen. Phys.Rev.Lett. 87, 147003 (2001)
[4] J.Schmalian, D.Pines, B.Stojkovic. Phys.Rev.Lett. 80, 3839(1998); Phys.Rev. B60, 667 (1999)
[5] E.Z.Kuchinskii, M.V.Sadovskii. Zh.Eksp.Teor.Fiz. 115, 1765 (1999); (JETP 88, 968 (1999)); ArXiv: cond-mat/9804129
[6] M.V.Sadovskii. Zh.Eksp.Teor.Fiz. 66, 1720 (1974); Fiz.Tverd.Tela 16, 2504 (1974)
[7] M.V.Sadovskii. Zh.Eksp.Teor.Fiz. 77, 2070(1979)
[8] A.I.Posazhennikova, M.V.Sadovskii. Zh.Eksp.Teor.Fiz. 115, 632 (1999); (JETP 88, 347 (1999)); ArXiv: cond-mat/9806199
[9] E.Z.Kuchinskii, M.V.Sadovskii. Zh.Eksp.Teor.Fiz. 117, 613 (2000); (JETP 90, 535 (2000)); ArXiv: cond-mat/9910261
[10] E.Z.Kuchinskii, M.V.Sadovskii. Zh.Eksp.Teor.Fiz. 119, 553 (2001); (JETP 92, 480 (2001)); ArXiv: cond-mat/0008377
[11] E.Z.Kuchinskii, M.V.Sadovskii. Zh.Eksp.Teor.Fiz. 121, 758 (2002); (JETP 94, No.3 (2002)); ArXiv: cond-mat/0110013
[12] M.V.Sadovskii, A.A.Timofeev. Superconductivity: Physics, Chemistry, Technology 4, 11(1991)
[13] M.V.Sadovskii, A.A.Timofeev. J.Moscow Phys.Soc. 1, 391 (1991)
[14] M.V.Sadovskii. Pis‘ma Zh.Eksp.Teor.Fiz. 69, 447 (1999); (JETP Lett. 69, 483 (1999)); Physica C341-348, 939 (2000); ArXiv: cond-mat/9902192
[15] P.Monthoux, A.Balatsky, D.Pines. Phys.Rev. B46, 14803 (1992)
[16] P.Monthoux, A.Balatsky. Phys.Rev. B47, 6069 (1993); Phys.Rev. B48, 4261 (1994)
[17] M.V.Sadovskii. Physica C341-348, 811 (2000); ArXiv: cond-mat/9912318
[18] L.Bartosch, P.Kopietz. Phys.Rev. B60, 15488 (1999)
[19] D.Vollhardt, P.Wolfle. Phys.Rev. B22, 4666 (1980)
[20] L.P.Gor’kov, A.I.Larkin, D.E.Khmelnitskii. Pis‘ma Zh.Eksp.Teor.Fiz. (JETP Lett.) 30,248 (1979)
[21] P.Monthoux, D.J.Scalapino. Phys.Rev. B65, 235104 (2002); ArXiv:cond-mat/0111534
[22] A.A.Gogolin, G.T.Zimanyi. Sol.St.Comm. 46, 469 (1983)
[23] X.J.Zhou, P.Bogdanov, S.A.Kellar, T.Noda, H.Eisaki, S.Uchida, Z.Hussain, Z.-X.Shen. Science 286, 268 (1999)
[24] A.Damascelli, D.H.Lu, Z.-X.Shen. ArXiv:cond-mat/0107042
[25] D.N.Basov, A.V.Puchkov, R.A.Hughes, T.Strach, J.Preston, T.Timusk. Phys.Rev. B49, 12165 (1994)
[26] D.N.Basov, B.Dabrowski, T.Timusk. Phys.Rev.Lett. 81, 2132 (1998)
[27] E.J.Singley, D.N.Basov, K.Kurahashi, T.Uefuji, K.Yamada. ArXiv:cond-mat/0103480
[28] Y.Onose, Y.Taguchi, K.Ishizaka, Y.Tokura. Phys.Rev.Lett. 87, 317001 (2001)