Hall number, optical sum rule and carrier density for the \( t\!-\!t'\!-\!J \) model

Jan O. Haerter and B. Sriram Shastry

Physics Department, University of California, Santa Cruz, California 95064, USA

(Received 13 August 2007; published 23 January 2008)

We revisit the relationship between three classical measures of particle number, namely, the chemical doping \( x \), the Hall number \( x_{\text{Hall}} \), and the particle number inferred from the optical sum rule \( x_{\text{opt}} \). We study the \( t\!-\!t'\!-\!J \) model of correlations on a square lattice, as a minimal model for high \( T_c \) systems, using numerical methods to evaluate the low temperature Kubo conductivities. These measures disagree significantly in this type of system, owing to Mott Hubbard correlations. The Hall constant has a complex behavior with several changes of sign as a function of filling \( x \), depending on the model parameters. Thus, \( x_{\text{Hall}} \) depends sensitively on \( t' \) and \( J \) due to a kind of quantum interference.

DOI: 10.1103/PhysRevB.77.045127

PACS number: 71.10.Fd, 72.15.Gd

I. INTRODUCTION

The traditional strategy, of converting a measured Hall constant or an optical sum rule to an electron count, runs into serious difficulties when interactions are strong within a lattice fermi system. The nonconservation of the lattice current, unlike its continuum counterpart, changes the \( f \)-sum rule drastically to involve nonuniversal variables such as the kinetic energy expectation. Similarly, the Hall constant suffers serious many body renormalization due to physics associated with the Mott Hubbard correlations; holes in the Mott insulator have little resemblance to carriers in uncorrelated bands.

This problem has very recently been revived in the context of \( \text{La}_2\!-\!\alpha\text{Sr}_\alpha\text{CuO}_4 \) (LSCO) the authors refining the initial work of Takagi and co-workers\cite{2} using high quality thin films. This is a particularly suitable system since the doping can be tuned all the way from the lightly doped to the overdoped Fermi liquid regime. The Hall constant provides several outstanding puzzles, first a change of sign from \( R_H>0 \) at small \( x \) to \( R_H<0 \) at \( x \approx 0.3 \), where \( x=1-n \) is the number of holes per copper. Further, there is a quite substantial \( T \) dependence for small \( x \approx 0.3 \). The problem is compounded by the angle resolved photoemission (ARPES) data,\cite{3} which shows that the topology of the Fermi surface remains electronlike from \( x \approx 0.18 \), so that the change of sign cannot be easily ascribed to a Fermi surface distortion. There is a notable recent attempt\cite{4} to rationalize the observed behavior using theoretical ideas\cite{5} invoking strong and anisotropic impurity (elastic) scattering. Thus, factors extrinsic to the two dimensional plane are invoked to understand the \( x, T \) dependence.

As noted recently,\cite{1} the measured Hall constant in better samples continues toward large negative values as \( x \rightarrow 1 \), in contrast to the early data\cite{2} that appeared to saturate. This overall behavior of the Hall constant, namely, a large positive value as \( x \rightarrow 0 \) and a large negative value as \( x \rightarrow 1 \) are precisely of the kind intrinsic to a Mott Hubbard system, as first pointed out in Ref. 6. Thus, a final theory would reconcile impurity scattering to intrinsic factors of the kind we study in this work. The early work of Ref. 6 [Shastry-Shraiman-Singh (SSS)] showed that the high frequency Hall constant shows a sensitivity to half filling and hence to Mott Hubbard physics. It gives a divergent Hall constant at half filling, together with (at least) three zero crossings, as the band filling \( n=\text{electrons}/\text{sites} \) varies from 0 to 2. Other recent ideas on Mott physics lead to comparable results. The results of SSS were obtained for a nearest neighbor \( t\!-\!J \) model on the square lattice at high temperature. This led to a holohlike Hall constant for \( 0 \leq x \leq 0.3 \), followed at large \( x \) by an electronic Hall constant. This change of sign is in agreement with the experiments\cite{1,2} on LSCO, but not so with several other High \( T_c \) compounds (e.g., \( \text{YBa}_2\!\text{Cu}_3\text{O}_{6+\delta} \)) that do not show a change of sign within the available range of doping. Thus, the problem of understanding the Hall constant in the various classes of high \( T_c \) systems remained unresolved, a task that we return to in this work.

Further recent experimental work of Refs. 8 and 9 on the optical mass and anomalous behavior of the Hall number in good samples of LSCO adds motivation to this effort. Here, we address the problem of computing the effect of correlations on the effective carrier count, or equivalently the Hall constant and the optical mass, for a model system, the \( t\!-\!t'\!-\!J \) model on a square lattice. While the optical mass is quite straightforward to address, using exact computation of the expectation value of the "stress tensor" or kinetic energy, the case of the Hall constant is quite non trivial, as elaborated below.

A study of the Fermi surface is another possible source of information on the Hall constant. We have alluded to the recent work in Ref. 3 on the ARPES derived shape of the Fermi surface for LSCO at all dopings. Theoretically, however, this is a vexed issue. First, in an interesting numerical study, the Luttinger theorem’s validity in \( t\!-\!J \) models describing strongly correlated matter has recently been questioned.\cite{10} Even when the theorem does apply, the possibility of shape deformation\cite{11} is strong. The implications are that for any choice of bare band parameters \( t, t' \) made, leading to anisotropic bare Fermi surfaces, one must exercise caution in interpreting the observed Fermi surface. This is so, since the Fermi surface is further deformed in an area (volume) preserving fashion due to the interactions, leading to the experimentally observed renormalized Fermi surface. The final observed Fermi surface is expected to be quite different from the starting shape since there are reasons to expect a strongly momentum dependent self-energy.\cite{12} There have been few studies of this difficult issue in literature since it requires the
knowledge of the momentum dependence of the self-energy. The situation for numerical studies is also rather unfavorable, since very few values of the momentum are available in finite sized clusters, making it hard to determine a surface. We therefore avoid any discussion of it and continue with a study of objects that are more direct for the Hall constant.

Motivated by the triangular lattice system Na$_x$CoO$_2$, we have recently studied the Hall constant as a function of temperature as well as frequency quite thoroughly. We used exact diagonalization to compute the exact Kubo formulas for the Hall constant to benchmark the high frequency approximations to the same. This parallel experience is helpful in the present context. Quite encouraging is the result that the frequency dependence is mild in essentially all cases studied, so that one can get a reasonable estimate of the Hall constant from the high frequency results. The temperature dependence of the Hall constant is serious for the triangular lattice, owing to the peculiar structure of the closed loops on the former. For the square lattice, this is not expected to be as serious, on general grounds. It is, however, found that the underdoped cases do have an inexplicable $T$ sensitivity, especially at low $T$ and low $x$, although the scale of the $T$ dependence is modest in comparison to that in the triangular lattice cobaltates. We are unable to address this issue here, as it seems to be related to the essential complexity of the pseudogap phase. Our computations are all at a low (effectively zero) temperature.

In this work, we go beyond the framework of SSS by studying the $t$-$t'$-$J$ model [Eq. (4)] on the square lattice (without the restriction of high-temperature expansions). This is an often used model to describe the physics of the copper oxide planes in high $T_c$ cuprates. The addition of the second neighbor hopping $t'$ is required by local density approximation calculations in order to fit to an effective tight binding model. Most importantly for our purpose, it extends nontrivially the simple $t$-$J$ model studied earlier and yields a rich variety of behavior of the Hall constant that seems to have the potential to explain the observed experimental diversity. In this work, we present preliminary results in this direction by computing the Hall constant on small clusters of the above model for various values of the ratio $t'/|t|$ for small clusters of up to 15 sites. We demarcate regions where the change of sign is observed as in SSS, from those where apparently no change occurs. Moreover, we provide rough estimates of the effective number of holes as a function of the chemical doping and the ratio $t'/|t|$. We are unable to examine more subtle issues such as the possible existence of a quantum critical point in Ref. 9, but rather wish to provide a rough base line from which one can build a more elaborate theory.

Given a theoretical model with $x$ holes per copper, one can compute an effective hopping $x_{Hall}$ from the Hall constant $R_H$ via $x_{Hall}=|v/(R_Hq_c)|$, where $v$ is the volume per copper and $q_c=|e|$ the elementary unit of charge. Similarly, given the optical conductivity $\sigma(\omega)$, we can define an optical hopping $x_{opt}$. Consider the $f$-sum rule on a lattice,

$$\tau^{xx} = \frac{\hbar^2}{2m_b} \sum_{k,\alpha} \frac{d^2 \varepsilon(k)}{dk^2} c_{\alpha}^*(k)c_{\alpha}(k)$$

the stress tensor. We can define the effective plasma frequency from $\omega_p = \sqrt{\frac{e^2 n}{m_b^*}}$, so that the $f$-sum rule leads to $\omega_p^2/8$ as usual. In the case of a parabolic noninteracting band, this object reduces to the familiar result and provides a natural generalization to the tight binding cases.

We note that the optical sum rule can also be interpreted as a renormalization of the effective mass since it only measures the ratio of filling to mass. We favor the above factorization, wherein the $x_{opt}$ contains all the many body renormalizations, but not the band effects. The band effects can be absorbed into the (optical) band mass $m_b$ meaningfully as follows. We define $m_b = m_b(t',x)$, where

$$\frac{\hbar^2 n}{m_b v} = \frac{1}{L} \sum_{k,\alpha} \frac{d^2 \varepsilon(k)}{dk^2} \langle c_{\alpha}(k)c_{\alpha}(k) \rangle_0,$$

with $n=1-x$ the electron density per copper with the average being carried out in the noninteracting band. The ratio $m_b/m_c$ as a function of its various arguments is easily evaluated, where $m_c$ is the bare electronic mass. For the case of $t'=0$ and $t=5160$ K, the ratio $m_b/m_c \approx 1.0$. In view of this close proximity between the band and bare masses, we simply set $m_b=m_c$. The lattice parameter used in our computations is $a_0=3.79 \times 10^{-10}$ m appropriate to LSCO, so the use of our results for other materials would require a small adjustment factor for the atomic volume.

In comparing with experiments, it must also be borne in mind that the projected $t$-$J$ model contains only a part of the spectral weight, since it describes the low energy part of the Hilbert space. Literally, it implies that the charge transfer gap is sent to infinity, so the integration in the sum rule must be cut off at roughly some fraction of the charge transfer gap. In practice, the upper limit for the frequency integral is often chosen precisely in such a way so that a comparison is not unjustified.

As stated above, in a weakly correlated system, all three particle numbers are expected to be equal; hence, $x=x_{Hall}=x_{opt}$. In strongly correlated systems, however, it is expected that this simple relation no longer holds since different variables undergo different renormalizations. Further, these many body effects also depend on the initial starting model parameters nontrivially, including the band structure effects. For the band structure in cuprate materials, several groups have emphasized the need to include second and possible further neighbor hoppings. In the following, we attempt to shed light on the different many body effects for a given chemical doping and for different “band parameters” $t$, $t'$ as well as $J$.

In Sec. II, we state the model and state the formulas that are computed as well as some indication of the methods used. In Sec. III, we discuss the results for the Hall constant, its frequency dependence, the effective Hall number, and the optical mass. In Sec. IV, we make concluding remarks.
II. MODEL AND EXACT DIAGONALIZATION

We study the $t$-$t'$-$J$ model on the square lattice, as a model for the strongly correlated cuprates.

\[
\hat{H} = -t \sum_{\langle i,j \rangle, \sigma} \hat{P}_G \hat{\epsilon}^\dagger_{i\sigma} \hat{c}^\dagger_{j\sigma} \hat{P}_G + t' \sum_{\langle\langle i,j \rangle \rangle, \sigma} \hat{P}_G \hat{\epsilon}^\dagger_{i\sigma} \hat{c}^\dagger_{j\sigma} \hat{P}_G
\]

\[
+ J \sum_{\langle i,j \rangle} \left( \hat{S}_i \cdot \hat{S}_j - \frac{\hat{n}_i \hat{n}_j}{4} \right),
\]

where $\hat{\epsilon}_{i\sigma}$ ($\hat{c}^\dagger_{i\sigma}$) creates (annihilates) an electron of spin $\sigma$, $\hat{S}_i$ is the three-component spin operator, $\hat{n}_i$ is the number operator, and $i$ specifies the lattice site. $\hat{P}_G$ denotes the Gutzwiller projector and the summation is over all nearest (second nearest) neighbor pairs $\langle i,j \rangle$ ($\langle\langle i,j \rangle \rangle$). Here, $t$ ($t'$) is the nearest (second nearest) neighbor hopping amplitude.

We have employed toroidal geometries with $L=14$ and 15 sites. Whenever possible, we reduce the computational effort by exploitation of space group symmetries. The symmetries are lowered upon the introduction of a magnetic field, as relevant for the evaluation of the Hall coefficient and the Hall number. For example, simple translations are no longer good symmetries. The magnetic field is introduced by the usual Peierls substitution which modifies the hopping $t$ between sites $i$ and $j$ by

\[
t \to t_{ij}(A) = t \exp \left( i \frac{2\pi}{\phi_0} \int_{i}^{j} A \cdot ds \right),
\]

where $A$ is the magnetic vector potential and $\phi_0 = \frac{hc}{e}$ is the flux quantum. We define the dimensionless flux threading a square plaquette as $\alpha = \frac{2\pi}{\phi_0} \oint \mathbf{A} \cdot ds$. In finite systems, the value of the smallest nonzero magnetic field is limited to values of $\alpha \geq \pi/1$, where $l$ is the length of a periodic loop in the system. Through a particular gauge we can achieve $l$ equal to the number of square faces in the cluster, this guarantees the equality of the flux values through all plaquettes. In the case of second neighbor hopping, we introduce additional phase factors along the diagonals of the square plaquettes in such a way that all fluxes through the resulting triangular plaquettes become equal. An example is given in Fig. 1. A similar strategy has been followed in the case of the square lattice quantum Hall effect. The Hall coefficient has been investigated earlier within the nearest-neighbor Hubbard and $t$-$J$ models. In this work, we are most interested in the dependence of this quantity on a second-neighbor hopping parameter. This task seems necessary to include in the starting model to explain the wide variety of behavior observed in the cuprates, and has not apparently been undertaken earlier.

III. RESULTS

A. Frequency-dependence of $R_H$

To further establish the validity of the high-frequency limit $R_H^\infty$ of the Hall coefficient we compute explicitly the real and imaginary parts of the frequency dependence of $R_H(\omega)$ through the Kubo formula, for the electrical conductivity $\sigma_{\alpha\beta}$

\[
\sigma_{\alpha\beta}(\omega) = \frac{i}{\omega \Omega} \left[ \langle \hat{\rho} \hat{\rho}^\dagger \rangle - \frac{1}{Z} \sum_{\nu} e^{-\beta \epsilon_{\nu}} - e^{-\beta \epsilon_{\mu}} \langle \nu | J_\alpha | \mu \rangle \right] \times \langle \mu | J_\beta | \nu \rangle,
\]

where $\beta$ is the inverse temperature, $\Omega$ is the volume of the system, $\hbar = 1$, and the sum is taken over all eigenstates of the system. The symbol $J_\alpha$ stands for the current operator in a field, and $Z$ is the partition function. The complex frequency dependent Hall coefficient can then be expressed as

\[
R_H(\omega) = \frac{i}{\omega \Omega} \left( \langle \hat{\rho} \hat{\rho}^\dagger \rangle - \frac{1}{Z} \sum_{\nu} e^{-\beta \epsilon_{\nu}} - e^{-\beta \epsilon_{\mu}} \langle \nu | J_\alpha | \mu \rangle \right) \times \langle \mu | J_\beta | \nu \rangle,
\]
We plan to return to more extensive computations in the future in order to extract the transport Hall constant. For the present computation of the Hall number $x_{\text{Hall}}$, with the above cautionary remark, we use the high frequency object,\(^6\)

\[
R_H^* = \lim_{B \to 0} -\frac{i\Omega}{Bq_e^2} \frac{\langle J_x J_y \rangle}{\langle \sigma^2 \rangle^2},
\]

(10)

**B. Hall coefficient**

We now analyze the doping dependence of the ground state Hall coefficient $R_H^*$ when a second neighbor hopping is included in the Hamiltonian of Eq. (4). We begin with $J=0$ (bottom panel of Fig. 3). We find that the value of a zero crossing at finite doping is in fact highly sensitive to the value of $t'$ (Fig. 3). At $t'=0$, the computations show a zero crossing near $x=1/3$ similar to the prediction from the high-temperature expansion.\(^6\) Turning on a positive $t'$, the zero crossing is pushed to lower $x$ and is essentially invisible in our studies since we cannot reach appreciably below $x=0.12$. Turning on a negative $t'$, the zero crossing is more pronounced and is pushed out to larger $x$. In order to place these results in context, recall that a positive $t'$ for hole doping leads to electronic frustration, as in the triangular lattice sodium cobaltate.\(^{13}\) A negative $t'$, on the one hand, causes a ferromagnetic Nagaoka tendency (toward a large Fermi surface). While quantum fluctuations as well as the pernicious influence of the exchange constant $J$ prevent the collapse into ordered states, these tendencies do seem to influence the behavior of the Hall constant. We thus interpret the strong dependence on the sign of $t'$ as a quantum interference effect. In our earlier study on the triangular lattice,\(^{13}\) we found results that are very similar to what we find here for $t'>0$.

To study the effect of $J>0$, we compute the Hall constant at two representative values of $J$ (two upper panels of Fig. 3). We see that exchange has a similar effect to $t'>0$, both lead to a suppression of the magnitude of the Hall constant. The influence on the zero crossing is more complex: in some cases, it is suppressed (in our computationally available range of $x$); in others, we find an extra zero crossing at lower $x$ where $R_H$ becomes negative. Presumably, at very small $x$, it turns positive and diverges due to the Mott Hubbard gap. This would imply that the Hall constant has a total of *three*
HALL NUMBER, OPTICAL SUM RULE AND CARRIER

PHYSICAL REVIEW B 77, 045127 (2008)

FIG. 3. (Color online) Hall coefficient in experimental units as a function of doping for different values of $t'$. The dots are from the experimental data in Ref. 1, extrapolated to $T \to 0$. In the top panel, $R_H^{\infty}(T=\infty)$ is the high-temperature limit for the case $t'=0$ (Ref. 6). In these curves, $J$ is varied from 0 to 0.4|$t'$. The bottom and top most sets of curves correspond to $-0.3 \leq |t'| \leq 0.3$ in steps of 0.1, whereas the middle one has $-0.5 \leq |t'| \leq 0$. The upper curves suggest that the number of zero crossings of the Hall constant is 3 for $0 \leq x \leq 1$, since at small enough $x$, $R_H$ must show an upturn toward $+\infty$, due to the Mott Hubbard insulating state at half-filling.

FIG. 4. (Color online) $t'$-$J$ dependence of the (dimensionless) $\langle \Psi_0| [J_x, J_y]| \Psi_0 \rangle$: red curves (green curves) correspond to $J=0$ ($J = 0.4|t'|$). Note that for $t'/|t| > 0$ a finite value of $J$ has little effect, while for the opposite sign the effect is very pronounced. This is due to the observation that a positive $t'/|t|$ plays the same role as a positive $J$ from the counter Nagaoka Thouless physics (Ref. 24).

zero crossings in the range $0 \leq x \leq 1$ (or six in the range $0 \leq x \leq 2$), in contrast to a single crossing for the uncorrelated case.

To gain a further understanding of the effect of $t'$, we introduce a finite value of $J=0.4|t|$ and compare with the case of $J=0$. In Fig. 4, we show for several values of $t'$ the numerator of the high-frequency Hall coefficient $R_H^{\infty}$. It is sufficient for our purposes to investigate this quantity, as it determines the possible zero crossings of the Hall coefficient. The denominator contains $\langle t' \rangle$, which is a rather well-behaved quantity, varying only slightly in magnitude by introduction of a small second-neighbor hopping. It vanishes in the limits of $x \to 0$ and $x \to 1$ and ultimately leads to a divergence of $R_H^{\infty}$ in these two limiting cases.

In Fig. 4, it is instructive to begin with the case of $t'=0$ (lowest panel). Here, for $J=0$, we obtain the zero crossing at $x=1/3$, similar to a prediction from high-temperature expansions. Introducing a finite $J$ shifts the zero crossing to lower dopings ($x \sim 0.15$). Here, $J$ acts as a source of antiferromagnetic correlations. Phenomenologically speaking, these antiferromagnetic correlations tend to resist a zero crossing. In a sense, this is similar to the effect of the triangular lattice with a frustrated hopping amplitude, where the zero crossing is shifted to lower dopings. If the second-neighbor hop $t' > 0$ of the sign corresponding to an electronically frustrated system is now explicitly introduced (left five panels of the figure), we find an almost perfect alignment of the two curves of different $J$. Thus, adding $t' > 0$ has a similar effect to adding an antiferromagnetic $J$. This is quite consistent with our premise that this effect can be interpreted in terms
is employed and its dependence on the sign of the hopping band limit. Filling where its relevance is much stronger than close to then dramatically destroys this state, especially near half-Thouless tendency toward a large Fermi surface. Finite may loosely attribute this to the ferromagnetic Nagaoka Thouless physics at play in frustrated triangular loops. On the other hand, if \( t' < 0 \), i.e., a nonfrustrated sign is employed (right five panels of figure), the divergence between the two curves becomes much more pronounced. We may loosely attribute this to the ferromagnetic Nagaoka Thouless tendency toward a large Fermi surface. Finite \( t \) then dramatically destroys this state, especially near half-filling where its relevance is much stronger than close to the band limit.

Thus, an understanding of the sign of the Hall constant and its dependence on the sign of the hopping \( t' \) seems to be closely linked to understanding the magnetic implications of the sign of \( t' \). We emphasize that these trends refer to the tendencies of the correlated matter toward various kinds of magnetically ordered states, but do not invoke any actual broken symmetries. Hence, these are a statement about underlying short ranged correlations in the many body system.

C. Optical sum rule and Hall number

To evaluate the optical sum, we make use of the relations Eqs. (1) and (3). This is shown in Fig. 5 for two values of \( J \). We observe that this quantity is strikingly different from the inverse Hall constant. The optics derived \( x_{opt} \) follows roughly the chemical doping \( x \) and increases in magnitude as a function of \( t' \). One noticeable feature is that a naive linear extrapolation of the small \( x \) results misses the origin slightly: thus presumably there is a change in slope for smaller \( x \approx 0.12 \). It shows a maximum at intermediate dopings \( x =0.6 \) as the trade-off for the stress tensor between the available hole and electron carriers is optimized here. In particular, \( x_{opt} \) remains unaffected by the change in sign of the Hall constant when it occurs. We now examine optimum doping, motivated by recent experimental results on the Hall number in this range of doping. Experimentally, the Hall number shows rather unusual nonlinear dependence on chemical doping \( x \). To understand this behavior, we examine more closely the high-frequency limit \( R_H \) near doping \( x=0.15 \). This corresponds to the introduction of two holes into finite systems of 14 and 15 sites. The case of a single hole is numerically ill-behaved. Hence, for \( x=2/L \), we study the dependence of \( R_H^* \) on \( t'/|t| \) and \( J \) in a physically meaningful range of values. In Fig. 6, we present the numerator of Eq. (10) as a function of \( t'/|t| \) for several values of \( J \) and the two systems studied. The figure shows that the dependence on \( t'/|t| \) is rather pronounced, leading to a zero crossing in the case of \( J=0 \). However, a small but finite value of \( J \) tends to destroy the strong \( t' \) dependence.

The particle number \( x_{opt} \) obtained from the optical sum rule is shown in Fig. 7. Its \( t'/|t| \) dependence is much weaker than that of \( \langle J_x J_y \rangle \). By combining Fig. 6 and 7, we obtain the inverse Hall number, shown in Fig. 8. The comparison of the \( t'/|t| \) and \( J \) dependences of this quantity with that of \( x_{opt} \) makes two points very clear: (i) The optical particle number and the Hall number are fundamentally different objects in strongly-correlated systems. (ii) The explanation of the experimentally measured nonlinear Hall number lies in a com-
that of the case \(x_{Hall}^{1}\) near optimal doping \(x=0.14\) (left) and \(x=0.13\) (right) computed on 14 (left) and 15 (right) site clusters. The red (green) curve is for \(J=0\) (\(J =0.4|t|\)). We see that \(x_{Hall}\) is a sensitive function of \(t'/|t|\) with a change of sign for \(J=0\) near \(t'/|t|=0.1\).

complicated interplay between the effect of finite—but probably small—\(J\) and a nonzero value of \(t'/|t|\) which allows an electronlike Hall coefficient in the optimally doped regime. Figure 8 shows the doping value of the zero crossing \(x_{zc}\) of \(R_H\) as a function of \(t'/|t|\) for two different values of \(J\) (compare Fig. 4). The data in this plot (upper panel) are for the hole-doped situation, and in the lower panel for the electron doped case using the transformation\(14\)

\[
R_H(t,-t',n) = -R_H(t,t',2-n).
\]

In the extreme limits \(|t'/|t|| \to \infty\), the value of \(x_{zc}\) approaches that of the case \(t'=0\) since these two cases correspond to nearest-neighbor hopping on a bipartite square lattice; hence, the sign is irrelevant in these limits. However, for \(0.6 \geq t'/|t| \geq 0.2\), the zero crossing appears to disappear in the ground state, within the limits of our calculation. This disappearance is rather independent of the value of \(J\).

\[
\text{FIG. 8. (Color online) } x_{Hall}^{1} \text{ as a function of } t'/|t| \text{ near optimal doping } x=0.14 \text{ (left) and } x=0.13 \text{ (right) computed on 14 (left) and 15 (right) site clusters. The red (green) curve is for } J=0 \text{ (} J =0.4|t|\text{). We see that } x_{Hall} \text{ is a sensitive function of } t'/|t| \text{ with a change of sign for } J=0 \text{ near } t'/|t|=0.1.\]

In the extreme limits \(|t'/|t|| \to \infty\), the value of \(x_{zc}\) approaches that of the case \(t'=0\) since these two cases correspond to nearest-neighbor hopping on a bipartite square lattice; hence, the sign is irrelevant in these limits. However, for \(0.6 \geq t'/|t| \geq 0.2\), the zero crossing appears to disappear in the ground state, within the limits of our calculation. This disappearance is rather independent of the value of \(J\).

\[
\text{FIG. 9. (Color online) The ground state Hall-coefficient zero-crossing doping } x_{zc} \text{ vs } t'/|t| \text{ for two different values of } J \text{ for hole doping (top panel) and electron doping (bottom panel), curves related by Eq. (11). Our finite systems do not allow determination of } x_{zc} \text{ for all } t'/|t|; \text{ we extrapolate doping dependence (Fig. 4) to obtain estimates for } x_{zc}.\]

IV. CONCLUSION

In the high \(T_c\) materials, spin fluctuation ideas by Kontani and co-workers,\(25\) lead to doping and frequency dependences of the Hall constant that are interesting. Also Kotliar and co-workers\(26\) have studied the Hall constant using dynamical mean field theory ideas. In this work, we have studied a strong coupling model, namely, the \(t-t'-J\) model by using a combination of theoretical ideas and computation of the exact spectrum of the model for small clusters. The problem addressed is that for classical metals the chemical doping \(x\), the Hall number \(x_{Hall}\), and particle number \(x_{opt}\) derived from the optical sum rule agree well. However, in strongly correlated systems, they follow completely different “renormalization paths.” In the present study of the \(t-t'-J\) model, we have extended previous studies to include the second neighbor hopping \(t'\). This term plays a crucial role in determining the detailed behavior of the Hall constant. The Hall number diverges at certain dopings and values of second neighbor hopping \(t'\). Furthermore, it strongly depends on the value of the interaction strength \(J\) for the case of positive (nonfrustrated) \(t'\). This unusual result is understandable in terms of the concept of “electronic frustration,” a form of quantum interference.

The inferred particle number \(x_{opt}\) increases roughly with the chemical doping \(x\) as the hole number increases. We do see a signature of a different slope for very small \(x \approx 0.12\). However, once the optimum trade-off between particle density and carrier freedom is reached, this quantity begins to decline and hence departs from the value of \(x\). Near optimal doping, we show that both \(t'\) and \(J\) significantly impact on the sign and magnitude of the Hall number. The optically derived hole number is much better behaved, i.e., its dependence on parameters is milder, and therefore seems a safer object to infer filling from.

In reconciling our numerical results with the recent experiments of Ref. 1 on clean films of LSCO, we concur with these authors that the data at \(T \approx 300\) K is safer to compare with the present type of theory since the \(T\) sensitivity is out of our theoretical reach. The absolute values of the Hall constant for \(x \approx 0.24\) found by them (their Fig. 3) are roughly comparable to what we find, although we do need to vary the parameters more systematically for attempting an actual fitting. Their recognition that larger \(x \approx 0.3\) leads to an unbounded growth of the (negative) Hall constant is important. It shows that the intrinsic behavior of data is in keeping with our ideas of Mott Hubbard physics versus uncorrelated band physics. This is explained in Ref. 1 and 6, where it is pointed out that in the limit \(x \to 1\), the Hall constant must be simply \(R_H \sim -v/q_j(1-x)\), due to the proximity of the band edge.

While our results are on quite small systems presently, they shed light on the questions arising from experiment, namely, a variety of changes of sign and unusual magnitudes of the Hall constant in different high \(T_c\) systems. This study also extends the insights of SSS (Ref. 6) and Stanescu and Phillips\(2\) on a Mott Hubbard theory of the Hall constant. Further detailed numerical studies could help produce systematic tables from which parameters could be inferred and thus help in subclassifying the high \(T_c\) materials more precisely.

045127-7
We gratefully acknowledge support from NSF Grant No. NSF-DMR0408247 and DOE-BES Grant No. DE-FG02-06ER46319. We thank F. Balakirev, D. Basov, G. Blumberg, G. H. Gweon, and H. Takagi for stimulating discussions.

ACKNOWLEDGMENTS

We gratefully acknowledge support from NSF Grant No. NSF-DMR0408247 and DOE-BES Grant No. DE-FG02-06ER46319. We thank F. Balakirev, D. Basov, G. Blumberg, G. H. Gweon, and H. Takagi for stimulating discussions.

10. The Fermi liquid relation (Ref. 11) \( m \frac{m}{m} = \left( \frac{\partial E(k, E(k))}{\partial k} \right) / \left( \frac{\partial E(k)}{\partial k} \right) \) implies that a vanishing of \( z_k \) leads to a divergent effective mass, unless the momentum dependence is divergent. Since the observed specific heat is not particularly enhanced in high \( T_c \) systems and further \( z_k \) is zero (or very small), we must conclude that the momentum dependence of the self-energy \( \Sigma(k, \omega) \) is very severe.